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FIELD OF THE INVENTION

present invention relates to novel recombinant which are mutants of naturally occurring allergens. Also, the invention relates to a composition comprising a mixture of the novel recombinant mutant allergens. Further, the invention relates to a method of preparing such recombinant mutant allergens as well as to pharmaceutical compositions, including vaccines, comprising the recombinant mutant allergens. In further embodiments, the present invention relates to methods of generating immune responses in a subject, vaccination or treatment of a subject as well as processes for preparing the compositions of the invention.

BACKGROUND OF THE INVENTION

Genetically predisposed individuals become sensitised 20 (allergic) to antigens originating from a variety of environmental sources, to the allergens of which the individuals are exposed. The allergic reaction occurs when a previously sensitised individual is re-exposed to 25 the same or a homologous allergen. Allergic responses range from hay fever, rhinoconductivitis, rhinitis and asthma to systemic anaphylaxis and death in response to e.g. bee or hornet sting or insect bite. The reaction is immediate and can be caused by a variety of atopic allergens such as compounds originating from grasses, 30 trees, weeds, insects, food, drugs, chemicals perfumes.

However, the responses do not occur when an individual is exposed to an allergen for the first time. The initial adaptive response takes time and does usually not cause

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any symptoms. But when antibodies and T cells capable of reacting with the allergen have been produced, any subsequent exposure may provoke symptoms. Thus, allergic responses demonstrate that the immune response itself can cause significant pathological states, which may be life threatening.

antibodies involved in atopic allergy primarily to immunoglobulins of the IgE class. IgE binds to specific receptors on the surface of mast cells and basophils. Following complex formation of a specific allergen with IgE bound to mast cells, receptor crosslinking on the cell surface results in signalling through the receptors and the physiological response of the target cells. Degranulation results in the release of i.a. histamine, heparin, a chemotactic factor eosinophilic leukocytes, leukotrienes C4, D4 and E4, which cause prolonged constriction of the bronchial smooth muscle cells. The resulting effects may be systemic or local in nature.

The antibody-mediated hypersensitivity reactions can be divided into four classes, namely type I, type II, type III and type IV. Type I allergic reactions is the classic immediate hypersensitivity reaction occurring within seconds or minutes following antigen exposure. These symptoms are mediated by allergen specific IgE.

Commonly, allergic reactions are observed as a response to protein allergens present e.g. in pollens, house dust mites, animal hair and dandruff, venoms, and food products.

In order to reduce or eliminate allergic reactions, 35 carefully controlled and repeated administration of allergy vaccines is commonly used. Allergy vaccination is traditionally performed by parenteral, intranasal, or sublingual administration in increasing doses over a fairly long period of time, and results in desensitisation of the patient. The exact immunological mechanism is not known, but induced differences in the phenotype of allergen specific T cells is thought to be of particular importance.

Allergy vaccination

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The concept of vaccination is based on two fundamental characteristics of the immune system, namely specificity and memory. Vaccination will prime the immune system of the recipient, and upon repeated exposure to similar proteins the immune system will be in a position to respond more rigorously to the challenge of for example a microbial infection. Vaccines are mixtures of proteins intended to be used in vaccination for the purpose of generating such a protective immune response in the recipient. The protection will comprise only components present in the vaccine and homologous antigens.

types Compared to other of vaccination allergy vaccination is complicated by the existence of an ongoing allergic patients. immune response in This response is characterised by the presence of allergen specific IgE mediating the release of allergic symptoms upon exposure to allergens. Thus, allergy vaccination using allergens from natural sources has an inherent risk of side effects being in the utmost consequence life threatening to the patient.

Approaches to circumvent this problem may be divided in three categories. In practise measures from more than one category are often combined. First category of measures includes the administration of several small doses over

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prolonged time to reach a substantial accumulated dose. Second category of measures includes physical modification of the allergens by incorporation of the allergens into gel substances such as aluminium hydroxide. Aluminium hydroxide formulation has adjuvant effect and a depot effect of slow allergen release reducing the tissue concentration of active allergen components. Third category of measures include chemical modification of the allergens for the purpose of reducing allergenicity, i.e. IgE binding.

The detailed mechanism behind successful allergy vaccination remains controversial. It is, however, agreed that T cells play a key role in the overall regulation of immune responses. According to current consensus the relation between two extremes of T cell phenotypes, Th1 and Th2, determine the allergic status of an individual. Upon stimulation with allergen Th1 cells secrete interleukines dominated interferon-y by leading protective immunity and the individual is healthy. Th2 cells on the other hand secrete predominantly interleukin 4 and 5 leading to IgE synthesis and eosinophilia and the individual is allergic. In vitro studies have indicated the possibility of altering the responses of allergen specific T cells by challenge with allergen derived peptides containing relevant T cell epitopes. approaches to new allergy vaccines are therefore largely based on addressing the T cells, the aim being to silence the T cells (anergy induction) or to shift the response from the Th2 phenotype to the Th1 phenotype.

Antibody-binding epitopes (B-cell epitopes)

X-ray crystallographic analyses of F_{ab}-antigen complexes
has increased the understanding of antibody-binding epitopes. According to this type of analysis antibody-

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binding epitopes can be defined as a section of the surface of the antigen comprising atoms from 15-25 amino acid residues, which are within a distance from the atoms of the antibody enabling direct interaction. The affinity of the antigen-antibody interaction can not be predicted the enthalpy contributed by van der interactions, hydrogen bonds or ionic bonds, alone. The entropy associated with the almost complete expulsion of water molecules from the interface represent an energy contribution similar in size. This means that perfect fit between the contours of the interacting molecules is a underlying antigen-antibody principal factor affinity interactions.

97/30150 (ref. 1), a population of protein molecules is claimed, which protein molecules have a distribution of specific mutations in the amino acid sequence as compared to a parent protein. From the description, it appears that the invention is concerned with producing analogues which are modified as compared to the parent protein, but which are taken up, digested and presented to T cells in the same manner as the parent Thereby, (naturally occurring allergens). protein modified T cell response is obtained. Libraries of modified proteins are prepared using a technique denoted PM (Parsimonious Mutagenesis).

In WO 92/02621 (ref. 2), recombinant DNA molecules are described, which molecules comprise a DNA coding for a polypeptide having at least one epitope of an allergen of trees of the order Fagales, the allergen being selected from $Aln\ g\ 1$, $Cor\ a\ 1$ and $Bet\ v\ 1$. The recombinant molecules described herein do all have an amino acid sequence or part of an amino acid sequence that corresponds to the sequence of a naturally occurring allergen.

WO 90/11293 (ref. 3) relates i.a. to isolated allergenic peptides of ragweed pollen and to modified ragweed pollen peptides. The peptides disclosed therein have an amino acid sequence corresponding either to the sequence of the naturally occurring allergen or to naturally occurring isoforms thereof.

Chemical modification of allergens

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Several approaches to chemical modification of allergens have been taken. Approaches of the early seventies include chemical coupling of allergens to polymers, and chemical cross-linking of allergens using formaldehyde, etc., producing the so-called 'allergoids'. The rationale behind these approaches was random destruction of IqE binding epitopes by attachment of the chemical ligand thereby reducing retaining IgE-binding while immunogenicity by the increased molecular weight of the complexes. Inherent disadvantages of 'allergoid' production are linked to difficulties in controlling the process of chemical cross-linking and difficulties in analysis and standardisation of the resulting high molecular weight complexes. 'Allergoids' are currently in clinical use and due to the random destruction of IqE binding epitopes higher doses can be administered as compared to conventional vaccines, but the safety and efficacy parameters are not improved over use of conventional vaccines.

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More recent approaches to chemical modification of allergens aim at a total disruption of the tertiary structure of the allergen thus eliminating IgE binding assuming that the essential therapeutic target is the allergen specific T cell. Such vaccines contain allergen sequence derived synthetic peptides representing minimal

T cells epitopes, longer peptides representing linked T cells epitopes, longer allergen sequence synthetic peptides representing regions of immunodominant T cell epitopes, or allergen molecules cut in two halves by recombinant technique. Another approach based on this rationale has been the proposal of the use of "low IgE binding" recombinant isoforms. In recent years it has become clear that natural allergens are heterogeneous containing isoallergens and variants having approximately 25% of their amino acids substituted. Some recombinant isoallergens have been found to be less efficient in IgE binding possibly due to irreversible denaturation and hence total disruption of tertiary structure.

In vitro mutagenesis and allergy vaccination

Attempts to reduce allergenicity by in vitro site directed mutagenesis have been performed using several allergens including Der f 2 (Takai et al, ref. 4), Der p 2 (Smith et al, ref. 5), a 39 kDa Dermatophagoides farinae allergen (Aki et al, ref. 6), bee venom phospholipase A2 (Förster et al, ref. 7), Ara h 1 (Burks et al, ref. 8), Ara h 2 (Stanley et al, ref. 9), Bet v 1 (Ferreira et al, ref. 10 and 11), birch profilin (Wiedemann et al, ref. 12), and Ory s 1 (Alvarez et al, ref. 13).

The rationale behind these approaches, again, is addressing allergen specific T cells while at the same time reducing the risk of IgE mediated side effects by reduction or elimination of IgE binding by disruption of the tertiary structure of the recombinant allergen. The rationale behind these approaches does not include the concept of dominant IgE binding epitopes and it does not include the concept of initiating a new

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protective immune response which also involves B-cells and antibody generation.

The article by Ferreira et al (ref. 11) describes the use of site directed mutagenesis for the purpose of reducing IgE binding. Although the three-dimensional structure of Bet v 1 is mentioned in the article the authors do not use the structure for prediction of solvent exposed amino acid residues for mutation, half of which have a low degree of solvent exposure. Rather they use a method developed for prediction of functional residues proteins different from the concept of structure based identification of conserved surface areas described here. Although the authors do discuss conservation of α -carbon backbone tertiary structure this concept is not a part of the therapeutic strategy but merely included to assess in vitro IqE binding. Furthermore, the evidence presented is not adequate since normalisation of CD-spectra prevents the evaluation of denaturation of a proportion of the sample, which is a common problem. The therapeutic strategy described aim at inducing tolerance in allergen specific T cells and initiation of a new immune response is not mentioned.

The article by Wiedemann et al. (ref. 12) describes the use of site directed mutagenesis and peptide synthesis the of monoclonal antibody purpose characterisation. The authors have knowledge of the tertiary structure of the antigen and they use this knowledge to select a surface exposed amino acid for mutation. The algorithm used can be said to be opposite to the one described by the present inventors since an acid differing from homologous sequences amino selected. The study demonstrates that substitution of a surface exposed amino acid has the capacity to modify the binding characteristics of a monoclonal antibody, which

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is not surprising considering common knowledge. The experiments described are not designed to assess modulation in the binding of polyclonal antibodies such as allergic patients' serum IgE. One of the experiments contained do apply serum IgE and although this experiment is not suitable for quantitative assessment, IgE binding does not seem to be affected by the mutations performed.

The article by Smith et al. (ref. 5) describes the use of site directed mutagenesis for the purpose of monoclonal antibody epitope mapping and reduction of IqE binding. The authors have no knowledge of the tertiary structure and make no attempt to assess the conservation of α carbon backbone tertiary structure. The algorithm used does not ensure that amino acids selected for mutation are actually exposed to the molecular surface. Only one of the mutants described lead to a substantial reduction in IgE binding. This mutant is deficient in binding of antibodies tested indicating that the structure is disrupted. The authors do not define a therapeutic strategy and initiation of a new immune response is not mentioned.

The article by Colombo et al. (ref. 14) describes the study of an IgE binding epitope by use of site directed mutagenesis and peptide synthesis. The authors use a three dimensional computer model structure based on the crystal structure of a homologous protein to illustrate the presence of the epitope on the molecular surface. The further presence of an epitope on a different allergen showing primary structure homology is addressed using synthetic peptides representing the epitope. The therapeutic strategy is based on treatment using this synthetic peptide representing a monovalent IqE binding epitope. Conserved surface areas between homologous allergens as well as the therapeutic concept

initiating a new protective immune response are not mentioned.

The article by Spangfort et al. (ref. 15) describes the three-dimensional structure and conserved surface exposed patches of the major birch allergen. The article does not mention major IgE binding epitopes nor site directed mutagenesis, neither is therapeutic application addressed.

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In none of the studies described above is IgE binding reduced by substitution of surface exposed amino acids while conserving α -carbon backbone tertiary structure. The rationale behind above-mentioned approaches does not include the concept of dominant IgE binding epitopes and it does not include the therapeutic concept of initiating a new protective immune response.

WO 99/47680 discloses the introduction of artificial amino acid substitutions into defined critical positions while retaining the α -carbon backbone tertiary structure of the allergen. In particular, WO 99/47680 discloses a recombinant allergen, which is a non-naturally occurring mutant derived from a naturally occurring allergen, wherein at least one surface-exposed, conserved amino acid residue of a B cell epitope is substituted by another residue which does not occur in the same position in the amino acid sequence of any known homologous protein within the taxonomic order from which said naturally occurring allergen originates, said mutant allergen having essentially the same α -carbon backbone tertiary structure as said naturally occurring allergen, and the specific IgE binding to the mutated allergen being reduced as compared to the binding naturally occurring allergen.

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The recombinant allergen disclosed in WO 99/47680 is obtainable by a) identifying amino acid residues in a naturally occurring allergen which are conserved with more than 70% identity in all known homologous proteins within the taxonomic order from which said naturally occurring allergen originates, b) defining at least one patch of conserved amino acid residues being coherently connected over at least 400 Å² of the surface of the three-dimensional of the allergen molecule as defined by having a solvent accessibility of at least 20%, said at least one patch comprising at least one B cell epitope, and c) substituting at least one amino acid residue in said at least one patch by another amino acid being nonconservative in the particular position while essentially preserving the overall α-carbon backbone structure of the allergen molecule.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1 shows mutant-specific oligonucleotide primers used for $Bet\ v\ 1$ mutant number 1. Mutated nucleotides are underlined.

Figure 2 shows two generally applicable primers (denoted "all-sense" and "all non-sense"), which were synthesised and used for all mutants.

Figure 3 shows the DNA and amino acid sequence of the naturally occurring allergen $Bet\ v\ 1$ as well as a number of $Bet\ v\ 1$ mutations.

Figure 4 shows the inhibition of the binding of biotinylated recombinant $Bet\ v\ 1$ to serum IgE from a pool of allergic patients by non-biotinylated $Bet\ v\ 1$ and by $Bet\ v\ 1$ Glu45Ser mutant.

shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 mutant Asn28Thr+Lys32Gln.

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inhibition of the shows the binding of biotinylated recombinant $Bet\ v\ 1$ to serum $IgE\ from\ a\ pool$ of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Pro108Gly mutant.

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Figure shows the inhibition of the binding biotinylated recombinant $Bet\ v\ 1$ to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Glu60Ser mutant.

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Figure 8 shows the CD spectra of recombinant and Triple-patch mutant, recorded at close equal concentrations.

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shows the inhibition of the binding biotinylated recombinant $Bet\ v\ 1$ to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Triple-patch mutant.

Figure 10 shows solvent accessibility of individually 25 aligned antigen 5 residues and alignment of Vespula antigen 5 sequences (left panel). On the right panel of Figure 10 is shown the molecular surface of antigen 5 with conserved areas among Vespula antigen 5:s.

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Figure 11 shows the sequence of the primer corresponding to the amino terminus of Ves v 5 derived from the sense strand. The sequence of the downstream primer is derived

from the non-sense strand.

Figure 12 shows two generally applicable primers (denoted

"all sense" and "all non-sense", which were synthesised and used for all mutants.

Figure 13 shows the DNA and amino acid sequence of the naturally occurring allergen $Ves\ v$ 5 as well as two $Ves\ v$ 5 mutations.

Figure 14 shows the inhibition of the binding of biotinylated recombinant Ves v 5 to serum IgE from a pool of allergic patients by non-biotinylated Ves v 5 and by Ves v 5 Lys72Ala mutant.

Figure 15 shows a theoretical model of the reaction between an allergen and mast cells by IgE cross-linking.

Figure 16 shows the DNA and amino acid sequence of the naturally occurring allergen $Der\ p$ 2.

Figure 17 shows schematically the primers used to create
20 the mutations. (I) shows the sense and antisense primers.
(II) shows the final recombinant protein harbouring
mutations at the indicated positions.

Figure 18 shows an illustration of the construction of
25 Bet v 1 mutants and a listing of the primers used. The
mutants contain from five to nine amino acids.

Figure 19 shows introduced point mutations at the surface of Bet v 1 (2628) and Bet v 1 (2637). In mutant Bet v 1 (2628), five primary mutations were introduced in one half of Bet v 1 leaving the other half unaltered. In mutant Bet v 1 (2637), five primary and three secondary mutations were introduced in the other half, leaving the first half unaltered.

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Figure 20 shows the circular dichroism (CD) spectra of

recombinant Bet v 1.2801 (wild type) and the Bet v 1 (2637) mutant recorded at nearly identical concentrations.

Figure 21 shows the inhibition of the binding of biotinylated recombinant Bet v 1.2801 (wild type) to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1.2801 and by Bet v 1 (2628), Bet v 1 (2637), and a 1:1 mix of Bet v 1 (2628) and Bet v1 (2637).

Figure 22 shows histamine release in human basophil cells of Bet v 1.2801 (wild type), Bet v 1 (2628), and Bet v 1 (2637).

Figure 23 shows histamine release in human basophil cells of Bet v 1.2801 (wild type), Bet v 1 (2628), and Bet v 1 (2637).

Figure 24 shows point mutations at the surface of Bet v 1 (2744).

Figure 25 shows point mutations at the surface of Bet v 1 (2753).

Figure 26 shows point mutations at the surface of Bet v 1 (2744) and Bet v 1 (2753).

Figur 27 shows circular dichroism (CD) spectra of Bet v 1.2801 (wild type) and Bet v 1 (2744), recorded at nearly equal concentrations.

Figur 28 shows histamine release in human basophil cells of Bet v 1.2801 (wild type), and mutant Bet v 1 (2744).

Figur 29 shows histamine release in human basophil cells

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of Bet v 1.2801 (wild type), and mutant Bet v 1 (2744).

Figur 30 shows point mutations at the surface of Bet v 1 (2733).

Figure 31 shows primers used for site-directed mutagenesis of Der p 2.

Figure 32 shows a sequence alignment of Der p 2 with other group 2 house dust mite allergens.

Figure 33 shows surface contours of Der p 2 from four different angles.

Figure 34 shows surface contours of a Der p 2 mutant from four different angles.

Figure 35A and B shows a sequence alignment of Der p 1 with other group 1 house dust mite allergens.

Figure 36 shows surface contours of Der p 1 from four different angles.

Figure 37 shows surface contours of a Der p 1 mutant from four different angles.

Figure 38A-D shows a sequence alignment of Phl p 5 with other group 5 grass allergens.

Figure 39A and B shows surface contours of Phl p 5 Model A and Model B, respectively, from four different angles.

Figure 40A and B shows surface contours of a Phl p 5 mutant Model A and B, respectively, from four different angles.

Figure 41 shows the proliferation of Peripheral Blood Lymphocytes expressed as Stimulation Index (SI) for various Bet v 1 preparations.

Figure 42-44 show the cytokine profile of T cells stimulated with various Bet v preparations. Figure 42 shows a patient with a Th0 profile, Figure 43 a Th1 profile and Figure 44 a Th2 profile.

10 OBJECT OF THE INVENTION

Rationale behind the present invention

The current invention is based on a unique rationale. According to this rationale the mechanism of successful allergy vaccination is not an alteration of the ongoing Th2-type immune response, but rather initiation of a new immune response involving tertiary epitope recognition by B-cells and antibody formation. It is believed that this new immune response is partly a Th1-type immune response. This model is supported by the observation that levels of specific IqE are unaffected by successful vaccination treatment, and that successful treatment is often accompanied by a substantial rise in allergen specific IgG4. In addition, studies of nasal biopsies before and after allergen challenge do not show a reduction in T cells with the Th2-like phenotype, but rather an increase in Th1-like T cells are observed. When vaccine the (or pharmaceutical compositions) administered through another route than the airways, it is hypothesised, that the new immune response evolves in a location physically separated from the ongoing Th2 response thereby enabling the two responses to exist in parallel.

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the assertion of the existence of so-called dominant IqE binding epitopes. It is proposed that these dominant IgE binding epitopes are constituted by tertiary structure surface dependent coherent areas large enough accommodate antibody binding conserved and isoallergens, variants, and/or homologous allergens from related species. The existence of cross-reactive IgE capable of binding similar epitopes on homologous allergens is supported by the clinical observation that allergic patients often react to several closely related species, e.g. alder, birch, and hazel, multiple grass species, or several species of the house dust mite genus Dermatophagoides. Ιt is furthermore supported laboratory experiments demonstrating IgE cross-reactivity between homologous allergens from related species and the capacity of one allergen to inhibit the binding of IgE to homologous allergens (Ipsen et al. 1992, ref. 16). It is well known that exposure and immune responses are related in a dose dependent fashion. Based on the combination of these observations it is hypothesised that conserved surface areas are exposed to the immune system in higher doses than non-conserved surface areas resulting in the generation of IgE antibodies with higher affinities, hence the term 'dominant IgE binding epitopes'.

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According to this rationale it is essential that the allergen has an α -carbon backbone tertiary structure which essentially is the same as that of the natural allergen, thus ensuring conservation of the surface topology of areas surrounding conserved patches representing targets for mutagenesis aimed at reducing IgE binding. By fulfilling these criteria the allergen has the potential to be administered in relatively higher doses improving its efficacy in generating a protective immune response without compromising safety.

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Furthermore, the invention is based on the finding that allergic symptoms are triggered by the cross-linking of allergen with two specific IgE's bound to the surface of effector cells, i.e. mast cells and basophils, via the high affinity IgE receptor, FceRI. For illustration, we refer to Fig. 15, which depicts a theoretical model of an allergen with IgE binding epitopes. Induction of mediator release from the mast cell and hence allergic symptoms is effected by allergen-mediated cross-linking of IgE bound to the surface of the mast cell, cf. Fig 15A. In the situation shown in Fig. 15B two of the epitopes have been mutated so as to reduce their IgE binding ability, and hence the allergen-mediated cross-linking is prevented. In this connection it should be noted that allergens usually comprise more than three B cell epitopes. However, from the theoretical situation depicted in Fig. 15 it may be assumed that the more epitopes, which have been mutated so as to eliminate or reduce their IqE binding ability, the lower the risk of allergen-mediated cross-linking and resulting allergic symptoms.

However, in order for a mutated allergen to be able to raise the new immune response, including the IgG response, the allergen must comprise at least one intact epitope. Preferably, the intact epitope is a dominant epitope, since such a mutated allergen will provide an improved protection when used for vaccination.

In conclusion, the inventive idea of the present invention is based on the recognition that a mutated 30 allergen having IgE binding reducing mutations multiple B cell epitopes, and at least one intact epitope, would on the one hand reduce the allergenmediated cross-linking and on the other hand allow the raising of 35 IgG response with a binding ability an competitive with that of IgE. Thus, the said mutated

allergen would constitute a highly advantageous allergen in that the risk of anaphylactic reactions would be strongly reduced.

Also, the present invention is based on the recognition that a vaccine comprising a mixture of different such mutated allergens, wherein ideally many or all epitopes are represented as intact, would be equally efficient in its ability to induce protection against allergic symptoms as the natural occurring allergen from which the mutated allergens are derived.

SUMMARY OF THE INVENTION

The present invention relates to the introduction of artificial amino acid substitutions into a number of defined critical positions, i.e. IgE binding epitopes, with the object of reducing the specific IgE binding capability of each mutated epitope.

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The invention provides а recombinant allergen, characterised in that it is a mutant of a naturally occurring allergen, wherein the mutant allergen has at least four primary mutations, which each reduce the specific IgE binding capability of the mutated allergen as compared to the IgE binding capability of the said naturally occurring allergen, wherein each primary mutation is a substitution of one surface-exposed amino acid residue with another residue, which does not occur in the same position in the amino acid sequence of any known homologous protein within the taxonomic species from which said naturally occurring allergen originates, wherein each primary mutation is spaced from each other primary mutation by at least 15 Å, and wherein the primary mutations are placed in such a manner that at least one circular surface region with a area of 800 ${\rm \AA}^2$

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comprises no mutation.

Without being bound by theory it is believed that the B cell epitopes can be distributed over almost the entire surface of the allergen. Furthermore, there experimental evidence that at least some epitopes constitute a part of a cluster of epitopes comprising a large number of overlapping epitopes. Therefore, the theoretical basis for the present invention is that any surface-exposed amino acid constitutes a potential site of mutation, which may result in a reduced capability to bind specific IqE.

Accordingly, the primary mutations are defined by their location in respect to each other, i.e. they are spaced apart, to ensure that they are mutations in separate clusters of epitopes.

The present invention also provides a composition comprising two or more recombinant mutant allergen variants according to claim 1, wherein each variant is defined by having at least one principal mutation, which is absent in at least one of the other variants, wherein for each variant no secondary mutation is present within a radius of 15 Å from each absent primary mutation. The composition preferably comprises 2-12, more preferably 3-10, more preferably 4-8 and most preferably 5-7 variants.

The present invention also provides a method of preparing the recombinant allergen according to claim 1, characterised in

a) identifying a number of amino acid residues in a naturally occurring allergen, which has a solvent accessibility of at least 20 %;

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b) selecting at least four of the identified amino acid residues in such a manner that each selected amino acid is spaced from each other selected amino acid by at least 15 Å, and that the selected amino acids are placed in such a manner that at least one circular surface region with a area of 800 Å 2 comprises no selected amino acid; and

- c) effecting for each of the selected amino acids a primary mutation, which reduce the specific IgE binding capability of the mutated allergen as compared to the binding capability of the said naturally occurring allergen, wherein each primary mutation is a substitution of a selected amino acid residue with another amino acid, which does not occur in the same position in the amino acid sequence of any known homologous protein within the taxonomic species from which said naturally occurring allergen originates.
- In an alternative aspect the invention relates to a method of preparing a recombinant allergen according to the invention, characterised in that the allergen is produced from a DNA sequence obtained by DNA shuffling (molecular breeding) of the DNA encoding the corresponding naturally occurring.

Furthermore, the invention relates to a recombinant allergen according to claim 1 for use as a pharmaceutical.

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Also, the invention relates to use of the recombinant allergen according to claim 1 for preparing a pharmaceutical for preventing and/or treating allergy.

Furthermore, the invention relates to the composition according to claim 37 for use as a pharmaceutical.

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Also, the invention relates to the use of a composition according to claim 37 for preparing a pharmaceutical for preventing and/or treating allergy.

the invention relates to a pharmaceutical composition, characterised in that it comprises recombinant allergen according to claim а composition according to claim 37, optionally in combination with a pharmaceutically acceptable carrier and/or excipient, and optionally adjuvant. an The pharmaceutical composition according to the invention may be in the form of a vaccine against allergic reactions

elicited by a naturally occurring allergen in patients

15 suffering from allergy.

Also, the invention relates to a method of generating an immune response in a subject comprising administering to the subject a recombinant allergen according to claim 1, a composition according to claim 37 or a pharmaceutical composition according to claim 41-42 or 46.

Further, the invention relates to vaccination or treatment of a subject comprising administering to the subject a recombinant allergen according to claim 1, a composition according to claim 37 or a pharmaceutical composition according to claim 41-42 or 46.

Also, the invention relates to a process for preparing a pharmaceutical composition according to claim 41 or 42 comprising mixing a recombinant allergen according to claim 1 or a composition according to claim 37 with pharmaceutically acceptable substances and/or excipients.

Further, the invention relates to a pharmaceutical composition obtainable by the process according to claim

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Also, the invention relates to a method for the treatment, prevention or alleviation of allergic reactions in a subject comprising administering to a subject a recombinant allergen according to claim 1, a composition according to claim 37 or a pharmaceutical composition according to claims 41-42 or 46.

Further, the invention relates to a DNA sequence encoding an allergen according to invention, a derivative thereof, a partial sequence thereof, a degenerated sequence thereof or a sequence, which hybridises thereto under stringent conditions, wherein said derivative, partial sequence, degenerated sequence or hybridising sequence encodes a peptide having at least one B cell epitope.

Also, the invention relates to an expression vector comprising the DNA according to the invention.

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Furthermore, the invention relates to a host cell comprising the expression vector according to the invention.

- 25 Additionally, the invention relates to a method of producing a recombinant mutant allergen comprising the step of cultivating the host cell according to the invention.
- Finally, the invention relates to a recombinant allergen according to the invention or encoded by the DNA sequence according to the invention comprising at least one T cell epitope capable of stimulating a T cell clone or T cell line specific for the naturally occurring allergen.
- The mutants according to invent should preferable be able to stimulate allergen specific T-cell lines in a similar

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manner/degree as measured by the T-cell stimulation index.

DETAILED DESCRIPTION OF THE INVENTION

In a preferred embodiment of the invention, the primary mutations are spaced 20 Å, preferably 25 Å and most preferably 30 Å.

10 It is believed that an allergen comprises a number of potential binding regions for specific IgE's, wherein each region approximately has a size of 800 Å², each surface region comprising a large number of overlapping epitopes. Thus, an allergen has a number of potential primary mutations of the surface area divided by 800 Å².

Preferably, the recombinant allergen according to the invention comprises from 5 to 20, preferably from 6 to 15, more preferably from 7 to 12, and most preferably from 8 to 10 primary mutations.

In a preferred embodiment of the invention, the surface region comprising no mutation has an area of 700 $\hbox{\AA}^2$, preferably 600 $\hbox{Å}^2$, more preferably 500 $\hbox{Å}^2$ and most preferably 400 $\hbox{Å}^2$.

preferred embodiment of the invention, recombinant allergen comprises a number of secondary mutations, which each reduce the specific IgE binding capability of the mutated allergen as compared to the binding capability of the said naturally occurring wherein each secondary mutation is substitution of one surface-exposed amino acid residue with another residue, which does not occur in the same position in the amino acid sequence of any known homologous protein within the taxonomic species from

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which said naturally occurring allergen originates, wherein the secondary mutations are placed outside the said circular region.

- The secondary mutations may be located close to a primary mutation, i.e. a secondary mutation may well be an additional mutation for the same epitope, which is mutated by the primary mutation.
- In a preferred embodiment of the invention, at least one of the surface-exposed amino acids to be substituted in the naturally occurring allergen has a solvent accessibility of above 20 %, preferably above 30 %, more preferably above 40 % and most preferably above 50 %.

In another preferred embodiment of the invention, at least one of the surface-exposed amino acids to be substituted in the naturally occurring allergen is conserved with more than 70 %, preferably 80 % and most preferably 90 % identity in all known homologous proteins within the species from which said naturally occurring allergen originates.

Preferably, the recombinant allergen according to invention essentially has the same α -carbon backbone tertiary structure as said naturally occurring allergen.

When comparing the α -carbon backbone tertiary structures of the mutant and the naturally occurring allergen molecules, the average root mean square deviation of the atomic coordinates is preferably below 2\AA .

In a preferred embodiment of the recombinant allergen of the invention, each amino acid residue to be incorporated into the mutant allergen does not occur in the same position in the amino acid sequence of any known

homologous protein within the taxonomic genus, preferably the subfamily, more preferably the family, more preferably the legion, more preferably the superfamily, more preferably the legion, more preferably the suborder and most preferably the order from which said naturally occurring allergen originates.

In a preferred embodiment of the invention the recombinant mutant allergen according to the invention is a non-naturally occurring allergen.

Specific IgE binding to the mutated allergen is preferably reduced by at least 5%, preferably at least 10% in comparison to naturally-occurring isoallergens or similar recombinant proteins in an immuno assay with sera from source-specific IgE reactive allergic patients or pools thereof.

Another way of assessing the reduced IgE binding and the reduced ability of mediating cross-linking of the mutant are the capability of the mutant to initiate Histamine Release (HR). The release of Histamine can be measured in several Histamine releasing assays. The reduced Histamine release of the mutants originates from reduced affinity toward the specific IgE bound to the cell surface as well as their reduced ability to facilitate cross-linking. HR is preferably reduced by 5-100%, more preferably 25-100%, more preferably 50-100% and most preferably 75-100% for the mutants of the invention in comparison to the naturally occurring allergens.

Typically, the circular surface region with an area of $800~\text{Å}^2$ comprising no mutation comprises atoms of 15-25 amino acid residues.

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A preferred recombinant allergen according to the

invention is characterised in that the surface-exposed amino acid residues are ranked with respect to solvent accessibility, and that one or more amino acids among the more solvent accessible ones are substituted.

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A further preferred recombinant allergen according to the invention is characterised in that the surface-exposed amino acid residues are ranked with respect to degree of conversation in all known homologous proteins within the species from which said naturally occurring allergen originates, and that one or more amino acids among the more conserved ones are substituted.

Preferably, the recombinant allergen according to the invention comprises from 1 to 4 secondary mutations per primary mutation.

A preferred embodiment of the invention is characterised in that one or more of the substitutions is carried out by site-directed mutagenesis.

Another preferred embodiment of the invention is characterised in that one or more of the substitutions is carried out by random mutagenesis.

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A further preferred embodiment of the invention is characterised in that one or more of the substitutions is carried out by DNA shuffling.

30 Recombinant allergens according to the invention may suitably be a mutant of an inhalation originating i.a. from trees, grasses, herbs, fungi, house dust mites, cockroaches and animal hair and dandruff. Important pollen allergens from trees, grasses and herbs 35 such originating from the taxonomic orders Fagales, Oleales and *Pinales* including i.a. birch

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(Alnus), alder hazel (Corylus), hornbeam and olive (Olea), the order of (Carpinus) Poales including i.a. grasses of the genera Lolium, Phleum, Poa, Cynodon, Dactylis and Secale, the orders of Asterales and Urticales including i.a. herbs of the genera Ambrosia and Artemisia. Important inhalation allergens from fungi are i.a. such originating from the genera Alternaria and Cladosporium. Other important inhalation allergens are from house dust mites of the Dermatophagoides, those from cockroaches and those from mammals such as cat, dog and horse. Further, recombinant allergens according to the invention may be mutants of venom allergens including such originating from stinging or biting insects such as those from the taxonomic order of Hymenoptera including bees (superfamily Apidae), wasps (superfamily Vespidea), and ants (superfamily Formicoidae).

Specific allergen components include e.g. Bet v 1 (B. verrucosa, birch), Aln g 1 (Alnus glutinosa, alder), Cor 20 a 1 (Corylus avelana, hazel) and Car b 1 (Carpinus betulus, hornbeam) of the Fagales order. Others are Cry j 1 (Pinales), Amb a 1 and 2, Art v 1 (Asterales), Par j 1 (Urticales), Ole e 1 (Oleales), Ave e 1, Cyn d 1, Dac q 25 1, Fes p 1, Hol l 1, Lol p 1 and 5, Pas n 1, Phl p 1 and 5, Poa p 1, 2 and 5, Sec c 1 and 5, and Sor h 1 (various grass pollens), Alt a 1 and Cla h 1 (fungi), Der f 1 and 2, Der p 1 and 2 (house dust mites, D. farinae and D. pteronyssinus, respectively), Lep đ 1 · and (Lepidoglyphus destructor; storage mite), Bla q 1 and 2, 30 Per a 1 (cockroaches, Blatella germanica and Periplaneta americana, respectively), Fel d 1 (cat), Can f 1 (doq), Equ c 1, 2 and 3 (horse), Apis m 1 and 2 (honeybee), Ves v 1, 2 and 5, Pol a 1, 2 and 5 (all wasps) and Sol i 1, 2, 3 and 4 (fire ant). 35

In one embodiment, the recombinant allergen is a mutant Bet v 1. Amino acids potentially suitable for substitution comprise amino acids V2, D72, E87, K-129, E-60, N-47, K-65, P-108, N-159, D-93, K-123, K-32, D-125, R-145, D-109, E-127, Q-36, E-131, L-152, E-6, E-96, D-156, P-63, H-76, E-8, K-134, E-45, T-10, V-12, K-20, S-155, H-126, P-50, N-78, K-119, V-2, L-24, E-42, N-4, A-153, I-44, E-138, G-61, A-130, R-70, N-28, P-35, S-149, K-103, Y-150, H-154, N-43, A-106, K-115, P-14, Y-5, K-137, E-141, E-87 and E-73. One or more of the primary and secondary substitutions may be selected from the group consisting of V2F, V2L, V2I, V2M, Y5V, T10P, T10A, K20N, D25E, N28T, K32Q, Q36A, Q36K, E42S, E45S, N47S, K55N, K65N, D72H, D72Q, D72N, T77A, N78K, E87G, E96L, K97S, K103V, P108G, D109N, K123I, D125Y, K129N, K134E, R145E, S149R, S149T, D156H and +160N, wherein + means that an additional amino acid is incorporated.

Examples of $Bet\ v\ 1$ mutants according to the present invention are as follows (parentheses, when used, indicate primary and secondary mutations):

Mutant A:

(Asn28Thr, Lys32Gln), (Asn78Lys, Lys103Val), Arg145Glu, (Asp156His, +160Asn).

Mutant B:

Tyr5Val, Glu42Ser, Glu45Ser, Asn78Lys, Lys103Val, Lys123Ile, Lys134Glu, Asp156His.

Mutant 2595 (Example 2): N28T, K32Q, E45S, P108G

Mutant 2628 (Example 4):

35 Tyr5Val, Glu45Ser, Lys65Asn, Lys97Ser, Lys134Glu.

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Mutant 2637 (Example 4):

Ala16Pro, (Asn28Thr, Lys32Gln), Lys103Thr, Pro108Gly, (Leu152Lys, Ala153Gly, Ser155Pro).

5 Mutant 2724:

N28T, K32Q, N78K, K103V, P108G, R145E, D156H, +160N.

Mutant 2733 (Example 4):

(Tyr5Val, Lys134Glu), (Asn28Thr, Lys32Gln), Glu45Ser,

Lys65Asn, (Asn78Lys, Lys103Val), Lys97Ser, Pro108Gly,

Arg145Glu, (Asp156His, +160Asn)

Mutant 2744: (Tyr5Val, Lys134Glu), (Glu42Ser, Glu45Ser), (Asn78Lys, Lys103Val), Lys123Ile, (Asp156His, +160Asn).

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Mutant 2753 (Example 4):

(Asn28Thr, Lys32Gln), Lys65Asn, (Glu96Leu, Lys97Ser), (Pro108Gly, Asp109Asn), (Asp125Tyr, Glu127Ser), Arg145Glu.

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Mutant 2744 + 2595:

Y5V, N28T, K32Q, E42S, E45S, N78K, K103V, P108G, K123I, K134E, D156H, +160N.

25 Mutant 2744 + 2628:

Y5V, E42S, E45S, K65N, N78K, K97S, K103V, K123I, K134E, D156H, +160N.

Mutant 2744 + 2595 + 2628:

30 Y5V, N28T, K32Q, E42S, E45S, K65N, N78K, K97S, K103V, P108G, K123I, K134E, D156H, +160N.

Furthermore, all of the above mutants comprising one or more of the following substitutions: V2F, V2L, V2I, V2M, T10A, K20N, Q36A or Q36K, D72H, D72Q, D72N, E87G, K129N

and S149R or S149T.

In another embodiment, the recombinant allergen is derived from a venom allergen from the taxonomic order of Vespidae, Apidae and Formicoidae.

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In a further embodiment, the recombinant allergen is derived from $Ves\ v$ 5. Amino acids potentially suitable substitution comprise amino acids Amino potentially suitable for substitution comprise amino acids K-16, K-185, K-11, K-44, K-210, R-63, K-13, F-6, K-149, K-128, E-184, K-112, F-157, E-3, K-29, N-203, N-34, K-78, K-151, L-15, L-158, Y-102, W-186, K-134, D-87, K-52, T-67, T-125, K-150, Y-40, Q-48, L-65, K-81, Q-101, Q-208, K-144, N-8, N-70, H-104, Q-45, K-137, K-159, E-205, N-82, A-111, D-131, K-24, V-36, N-7, M-138, T-209, V-84, K-172, V-19, D-56, P-73, G-33, T-106, N-170, L-28, T-43, Q-114, C-10, K-60, N-31, K-47, E-5, D-145, V-38, A-127, D-156, E-204, P-71, G-26, Y-129, D-141, F-201, R-68, N-200, D-49, S-153, K-35, S-39, Y-25, V-37, G-18, W-85 and more of the primary and I-182. One or secondary substitutions may be selected from the group consisting of K29A, T67A, K78A, V84S, Y102A, K112S, K144A, K202M and N203G.

In a further embodiment, the recombinant allergen is derived from Der p 2. Amino acids potentially suitable for substitution comprise amino acids R-128, D-129, H-11, H-30, S-1, K-77, Y-75, R-31, K-82, K-6, K-96, K-48, K-55, K-89, Q-85, W-92, I-97, H-22, V-65, S-24, H-74, K-126, L-30 61, P-26, N-93, D-64, I-28, K-14, K-100, E-62, I-127, E-102, E-25, P-66, L-17, G-60, P-95, E-53, V-81, K-51, N-103, Q-2, N-46, E-42, T-91, D-87, N-10, M-111, C-8, H-124, I-68, P-79, K-109 and R-128, D-129, H-11, H-30, S-1, K-77, Y-75, R-31, K-82, K-6, K-96, K-48, K-55, K-89, Q-85, W-92, I-97, H-22, V-65, S-24, H-74, K-126, L-61, P-26, N-93, D-64, I-28, K-14, K-100, E-62, I-127, E-102, E-

25, P-66, L-17, G-60, P-95, E-53, V-81, K-51, N-103, Q-2, N-46, E-42, T-91, D-87, N-10, M-111, C-8, H-124, I-68, P-79, K-109, K-15. One or more of the primary and secondary substitutions may be selected from the group consisting of K6A, N10S, K15E, S24N, H30N, K48A, E62S, H74N, K77N, K82N, K100N and R128O.

Examples of $Bet\ v\ 1$ mutants according to the present invention are as follows:

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Mutant A:

K6A, K15E, H30N, E62S.

Mutant B:

15 K6A, K15E, H30N, E62S, H74N, K82N.

Mutant C:

K6A, N10S, K15E, S24N, H30N, K48A, E62S, H74N, K77N, K82N, K100N and R128Q

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Vaccines

Preparation of vaccines is generally well known in the art. Vaccines are typically prepared as injectables either as liquid solutions or suspensions. Such vaccine may also be emulsified or formulated so as to enable nasal administration as well as oral, including buccal and sublingual, administration. The immunogenic component in question (the recombinant allergen as defined herein) may suitably be mixed with excipients which are pharmaceutically acceptable and further compatible with the active ingredient. Examples of suitable excipients are water, saline, dextrose, glycerol, ethanol and the like as well as combinations thereof. The vaccine may additionally contain other substances such as wetting agents, emulsifying agents, buffering agents or adjuvants enhancing the effectiveness of the vaccine.

Vaccines are most frequently administered parenterally by subcutaneous or intramuscular injection. Formulations which are suitable for administration by another route include oral formulations and suppositories. Vaccines for oral administration may suitably be formulated with excipients normally employed for such formulations, e.g. pharmaceutical grades of mannitol, lactose, magnesium sodium stearate, saccharine, cellulose, magnesium carbonate and the like. The composition can be formulated as solutions, suspensions, emulsions, tablets, pills, capsules, sustained release formulations, aerosols, powders, or granulates.

15 The vaccines are administered in a way so as to be compatible with the dosage formulation and in such amount as will be therapeutically effective and immunogenic. The quantity of active component contained within the vaccine depends on the subject to be treated, i.a. the capability of the subject's immune system to respond to the treatment, the route of administration and the age and weight of the subject. Suitable dosage ranges can vary within the range from about 0.0001 µg to 1000 µg.

As mentioned above, an increased effect may be obtained by adding adjuvants to the formulation. Examples of such adjuvants are aluminum hydroxide and phosphate (alum) or calcium phosphate as a 0.05 to 0.1 percent solution in phosphate buffered saline, synthetic polymers of sugars or polylactid glycolid (PLG) used as 0.25 percent solution. Mixture with bacterial cells such as C. parvum, endotoxins or lipopolysaccharide components of gramnegative bacteria, emulsion in physiologically acceptable oil vehicles such as mannide monoaleate (Aracel A) or emulsion with 20 percent solution of a perfluorocarbon (e.g. Fluosol-DA) used as a block substitute may also be

employed. Oil emulsions, such as MF-59 may also be used. Other adjuvants such as Freund's complete and incomplete adjuvants as well as saponins, such as QuilA, Qs-21 and ISCOM, and RIBI may also be used.

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Most often, multiple administrations of the vaccine will be necessary to ensure an effect. Frequently, the vaccine is administered as an initial administration followed by subsequent inoculations or other administrations. The number of vaccinations will typically be in the range of from 1 to 50, usually not exceeding 35 vaccinations. Vaccination will normally be performed from biweekly to bimonthly for a period of 3 months to 5 years. This is contemplated to give desired level of prophylactic or therapeutic effect.

The recombinant allergen may be used as a pharmaceutical preparation, which is suitable for providing a certain protection against allergic responses during the period of the year where symptoms occur (prophylaxis). Usually, the treatment will have to be repeated every year to maintain the protective effect. Preparations formulated for nasal, oral and sublingual application are particular suited for this purpose.

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Method of preparing a recombinant allergen according to the invention

As mentioned above, the present invention also relates to 30 a method of preparing the recombinant mutant allergen of the invention, cf. claim 48.

The surface-exposed amino acids suitable for substitution in accordance with the present invention may be identified on the basis of information of their solvent (water) accessibility, which expresses the extent of

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surface exposure. A preferred embodiment of the method of the invention is characterised in ranking the said identified amino acid residues with respect to solvent accessibility and substituting one or more amino acids among the more solvent accessible ones.

second parameter, which may contribute to the identification of surface-exposed amino acids suitable for substitution in accordance with the present invention is the extent in which an amino acid is conserved in all known homologous proteins within the species from which said naturally occurring allergen originates. Alternatively, the extent in which in all homologous proteins within the taxonomic subfamily, family, superfamily, legion suborder or order from which said naturally occurring allergen originates is used as such a second parameter.

Accordingly, a preferred embodiment of the method of the invention is characterised in selecting identified amino acid residues, which are conserved with more than 70 %, preferably more than 80 % and most preferably more than 90 % identity in all known homologous proteins within the species from which said naturally occurring allergen originates.

Furthermore, a particularly preferred embodiment of the method of the invention is characterised in ranking the said identified amino acid residues with respect to degree of conversation in all known homologous proteins within the species from which said naturally occurring allergen originates and substituting one or more amino acids among the more conserved ones.

35 A further preferred embodiment of the method of the invention comprises selecting the identified amino acids

so as to form a mutant allergen, which has essentially the same α -carbon backbone tertiary structure as said naturally occurring allergen.

- Another preferred embodiment of the method of the invention is characterised in that the substitution of amino acid residues is carried out by site-directed mutagenesis.
- An alternative preferred embodiment of the method of the invention is characterised in that the substitution of amino acid residues is carried out by DNA shuffling.

Criteria for substitution

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For molecules for which the tertiary structure has been determined (e.g. by x-ray crystallography, or NMR electron microscopy), the mutant carrying the substituted amino acid(s) should preferably fulfil the following criteria:

The overall α -carbon backbone tertiary structure of molecule is preferably conserved. Conserved defined as an average root mean square deviation of the atomic coordinates comparing the structures below 2Å. This is important for two reasons: a) It is anticipated that the entire surface of the natural constitutes an overlapping continuum of potential antibody-binding epitopes. The majority of the surface of the molecule is not affected by the substitution(s), and thus retain its antibody-binding inducing properties, which is important for the generation of new protective antibody specificities being directed at epitopes present also on the natural allergen. b) Stability, both concerning shelf-life and upon injection into fluids.

2. The amino acids to be substituted are preferably located at the surface, and thus accessible for antibody-binding. Amino acids located on the surface in the three-dimensional structure usually have a solvent (water) accessibility of at least 20%, suitably 20-80%, more suitably 30-80%. Solvent accessibility is defined as the area of the molecule accessible to a sphere with a radius comparable to a solvent (water, r = 1.4 Å) molecule.

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- Each of the substituted amino acids is preferably located in conserved patches having an area larger than 400 $Å^2$. Conserved patches are defined as coherently connected areas of surface-exposed conserved amino acid residues and backbone. Conserved amino acid residues are defined by sequence alignment of all known (deduced) amino acid sequences of homologues proteins within the taxonomic species, same genus, subfamily, family, superfamily, legion, suborder or order. Amino acid positions having identical amino acid residues in more than 70% of the sequences are considered conserved. Conserved patches are expected to contain epitopes to which the IgE of the majority of patients is directed.
- Conservation of α -carbon backbone tertiary structure is best determined by obtaining identical structures by x-ray crystallography or NMR before and after mutagenesis. In absence of structural data describing the mutant indistinguishable CD-spectra or immunochemical data, e.g. antibody reactivity, may render conservation of α -carbon backbone tertiary structure probable, if compared to the data obtained by analysis of a structurally determined molecule.
- 35 4. Within the conserved patches amino acids for mutagenesis should preferentially be selected among the

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most solvent (water) accessible ones located preferably near the centre of the conserved patch.

5. Preferentially, a polar amino acid residue is substituted by another polar residue, and a non-polar amino acid residue is substituted by another non-polar residue.

With an object of essentially retaining the three-dimensional structure of the allergen, the amino acid to be incorporated may be selected on the basis of a comparison with a protein, which is a structural homologue to the allergen, e.g. a protein, which belongs to the same taxonomic order as the allergen, and which does not have any cross-reactivity with the allergen.

DNA according to the invention

In a preferred embodiment, the DNA sequence of the invention is a derivative of the DNA sequence encoding the naturally occurring allergen.

Preferably, the DNA derivative is obtained by sitedirected or random mutagenesis of the DNA encoding the naturally occurring allergen.

In a first particularly preferred embodiment, the DNA sequence is a derivative of the sequence shown in Fig. 3, wherein the DNA sequence is mutated so as to encode an allergen having at least four mutations selected from the group consisting of K-129, E-60, N-47, K-65, P-108, N-159, D-93, K-123, K-32, D-125, R-145, D-109, E-127, Q-36, E-131, L-152, E-6, E-96, D-156, P-63, H-76, E-8, K-134, E-45, T-10, V-12, K-20, S-155, H-126, P-50, N-78, K-119, V-2, L-24, E-42, N-4, A-153, I-44, E-138, G-61, A-130, R-70, N-28, P-35, S-149, K-103, Y-150, H-154, N-43, A-106,

K-115, P-14, Y-5, K-137, E-141, E-87, E-73.

In a second particularly preferred embodiment, the DNA sequence is a derivative of the sequence shown in Fig. 13, wherein the DNA sequence is mutated so as to encode an allergen having at least four mutations selected from the group consisting of K-16, K-185, K-11, K-44, K-210, R-63, K-13, F-6, K-149, K-128, E-184, K-112, F-157, E-3, K-29, N-203, N-34, K-78, K-151, L-15, L-158, Y-102, W-186, K-134, D-87, K-52, T-67, T-125, K-150, Y-40, Q-48, 10 L-65, K-81, Q-101, Q-208, K-144, N-8, N-70, H-104, Q-45, K-137, K-159, E-205, N-82, A-111, D-131, K-24, V-36, N-7, M-138, T-209, V-84, K-172, V-19, D-56, P-73, G-33, T-106, N-170, L-28, T-43, Q-114, C-10, K-60, N-31, K-47, E-5, D-145, V-38, A-127, D-156, E-204, P-71, G-26, Y-129, D-141, 15 F-201, R-68, N-200, D-49, S-153, K-35, S-39, Y-25, V-37, G-18, W-85 and I-182.

In a third particularly preferred embodiment, the DNA 20 sequence is a derivative of the sequence shown in Fig. 16, wherein the DNA sequence is mutated so as to encode an allergen having at least four mutations selected from the group consisting of R-128, D-129, H-11, H-30, S-1, K-77, Y-75, R-31, K-82, K-6, K-96, K-48, K-55, K-89, Q-85, W-92, I-97, H-22, V-65, S-24, H-74, K-126, L-61, P-26, N-25 93, D-64, I-28, K-14, K-100, E-62, I-127, E-102, E-25, P-66, L-17, G-60, P-95, E-53, V-81, K-51, N-103, Q-2, N-46, E-42, T-91, D-87, N-10, M-111, C-8, H-124, I-68, P-79, K-109 and R-128, D-129, H-11, H-30, S-1, K-77, Y-75, R-31, K-82, K-6, K-96, K-48, K-55, K-89, Q-85, W-92, I-97, H-30 22, V-65, S-24, H-74, K-126, L-61, P-26, N-93, D-64, I-28, K-14, K-100, E-62, I-127, E-102, E-25, P-66, L-17, G-60, P-95, E-53, V-81, K-51, N-103, Q-2, N-46, E-42, T-91, D-87, N-10, M-111, C-8, H-124, I-68, P-79, K-109, K-15.

The recombinant mutant allergen according to the present invention may be produced using a DNA sequence obtained by DNA shuffling (molecular breeding) of the corresponding naturally DNA. DNA shuffling may be carried out according to the procedures disclosed in the article by Punnonen et al. (ref. 25) as well as the procedures disclosed in the articles mentioned therein, which are all included herein by this reference.

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Diagnostic assay

Furthermore, the recombinant mutant allergens according invention have diagnostic possibilities advantages. Prior art allergy vaccines are based on extracts of the naturally occurring allergen source, and thus represent a wide variety of isoforms. The allergic individual has initially been sensitised and has IgE to one or some of the isoforms present. Some of the isoforms may be relevant with respect to the allergic reactions of the allergic individual due to homology and subsequent cross-reactivity with the isoform to which the individual is allergic, whereas other isoforms may be irrelevant as they do not harbour any of the IqE binding epitopes to which the allergic individual has specific IqE. Due to this heterogeneity of the specificities of the population, some isoforms may therefore be safe to administer, i.e. they do not result in an allergic response via IgE, whereas other isoforms may be harmful causing undesirable side-effects.

Thus, the mutants of the invention and the compositions of the invention intended to be administered therapeutically may also be used for an in vivo or in vitro diagnostic assay to monitor the relevance, safety or outcome of a treatment with such mutants or

compositions. Diagnostic samples to be applied include body samples, such as sera.

Thus, the invention also relates to a diagnostic assay for assessing relevance, safety or outcome of therapy of a subject using a recombinant mutant allergen according to the invention or a composition according to the invention, wherein an IgE containing sample of the subject is mixed with said mutant or said composition and assessed for the level of reactivity between the IgE in said sample and said mutant. The assessing of the level of reactivity between the IgE in the sample and the mutant may be carried out using any known immunoassay.

15 Definitions

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In connection with the present invention the expression "reduce the specific IgE binding capability as compared to the IgE binding capability of the said natural-occurring allergen" means that the reduction is measurable in a statistically significant manner (p <0.05) in at least one immunoassay using serum from a subject allergic to the natural-occurring allergen. Preferably, the IgE binding capability is reduced by at least 5 %, more preferably at least 10 %.

The expression "surface-exposed amino acid" means that the amino acid residue is located at the surface of the three-dimensional structure in such a manner that when the allergen is in solution at least a part of at least one atom of the amino acid residue is accessible for contact with the surrounding solvent. Preferably, the amino acid residue in the three-dimensional structure has a solvent (water) accessibility of at least 20%, suitably at least 30 %, more suitably at least 40 % and most preferably at least 50 %.

The expression "solvent accessibility" is defined as the area of the molecule accessible to a sphere with a radius comparable to a solvent (water, r = 1.4 Å) molecule.

The expressions "surface-exposed" and "solvent-exposed" are used interchangeably.

The expression "the taxonomic species from which said naturally occurring allergen originates" means species in the taxonomic system.

Furthermore, the expression "said mutant allergen having essentially the same α -carbon backbone tertiary structure as said naturally occurring allergen" means that when comparing the structures, the average root mean square deviation of the atomic coordinates is below 2 Å.

In connection with the present invention the expression "substitution" means the deletion, substitution or addition of an amino acid in comparison to the amino acid sequence of the naturally occurring allergen.

The present invention is further illustrated by the following non-limiting examples.

EXAMPLES

EXAMPLE 1

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Example 1 describes the preparation of recombinant mutant allergens with one and three primary mutations. Recombinant mutant allergens according to the invention, i.e. allergens comprising at least four primary mutations, may be prepared using the same procedures.

Identification of common epitopes within Fagales pollen allergens

The major birch pollen allergen $Bet\ v\ 1$ shows about 90% amino acid sequence identity with major allergens from pollens of taxonomically related trees, i.e Fagales (for instance hazel and hornbeam) and birch pollen allergic patients often show clinical symptoms of allergic cross-reactivity towards these $Bet\ v\ 1$ homologous proteins.

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Bet v 1 also shows about 50-60% sequence identity with allergic proteins present in certain fruits (for instance apple and cherry) and vegetables (for instance celery and carrot) and there are clinical evidence for allergic cross-reactivity between $Bet\ v$ 1 and these food related proteins.

In addition, Bet v 1 shares significant sequence identity (20-40%) with a group of plant proteins called pathogenesis-related proteins (PR-10), however there are no reports of allergic cross-reactivity towards these PR-10 proteins.

Molecular modelling suggests that the structures of Fagales and food allergens and PR-10 proteins are close to being identical with the $Bet\ v\ 1$ structure.

The structural basis for allergic $Bet\ v\ 1$ cross-reactivity was reported in (Gajhede et al 1996, ref. 17) where three patches on the molecular surface of $Bet\ v\ 1$ could be identified to be common for the known major tree pollen allergens. Thus, any IgE recognising these patches on $Bet\ v\ 1$ would be able to cross-react and bind to other Fagales major pollen allergens and give rise to allergic symptoms. The identification of these common patches was performed after alignment of all known amino acid

sequences of the major tree pollen allergens in combination with an analysis of the molecular surface of Bet v 1 revealed by the α -carbon backbone tertiary structure reported in ref. 17. In addition, the patches were defined to have a certain minimum size (>400 Ų) based on the area covered by an antibody upon binding.

Selection of amino acid residues for site-directed mutagenesis

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Amino acid residues for site-directed mutagenesis were selected among residues present in *Bet v* 1 specific areas and the common patches since modifications of these is expected to affect the binding of serum IgE from the majority of patients showing clinical tree pollen allergic cross-reactivity.

The relative orientation and percentage of solvent-exposure of each amino acid residue within respective patch was calculated based on their atomic coordinates. Residues having a low degree of solvent exposure (<20%) were not regarded relevant for mutagenesis due to the possible disruption of the structure or lack of antibody interaction. The remaining residues were ranked according to their degree of solvent-exposure.

Sequence alignment

Sequences homologous to the query sequence (Bet v 1 No. 2801, WHO IUIS Nomenclature Subcommittee on Allergens) were derived from GenBank and EMBL sequence databases by a BLAST search (Altschul et al., ref. 18). All sequences with BLAST reported probabilities less than 0.1 were taken into consideration and one list were constructed containing a non-redundant list of homologous sequences. These were aligned by CLUSTAL W (Higgins et al., ref. 19)

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and the percentage identity were calculated for each position in the sequence considering the complete list or taxonomically related species only. A total of 122 sequences were homologous to $Bet\ v\ 1\ No.\ 2801$ of which 57 sequences originates from taxonomically related species.

Cloning of the gene encoding Bet v 1

RNA was prepared from Betula verrucosa pollen (Allergon, Sweden) by phenol extraction and LiCl precipitation. Oligo(dT)-cellulose affinity chromatography was performed batch-wise in Eppendorph tubes, and double-stranded cDNA was synthesised using a commercially available kit (Amersham). DNA encoding Bet v 1 was amplified by PCR and cloned. In brief, PCR was performed using cDNA as template, and primers designed to match the sequence of the cDNA in positions corresponding to the amino terminus of Bet v 1 and the 3'-untranslated region, respectively. The primers were extended in the 5'-ends to accommodate restriction sites (NcoI and HindIII) for directional cloning into pKK233-2.

Subcloning into pMAL-c

The gene encoding $Bet\ v\ 1$ was subsequently subcloned into 25 the maltose binding protein fusion vector pMAL-c (New England Biolabs). The gene was amplified by PCR and subcloned in frame with malE to generate maltose binding protein (MBP)-Bet v 1 protein fusion operons in which MBP 30 and $Bet\ v$ 1 were separated by a factor X_a protease clevage site positioned to restore the authentic aminoterminal sequence of Bet v 1 upon cleavage, as described in ref. 15. In brief, PCR was performed using pKK233-3 with $Bet\ v\ 1$ inserted as template and primers corresponding to the amino- and carboxyterminus of the 35 protein, respectively. The promoter proximal primer was

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extended in the 5'-end to accommodate 4 codons encoding an in frame factor X_a protease cleavage site. Both primers were furthermore extended in the 5'-ends to accommodate restriction sites (KpnI) for cloning. The Bet v 1 encoding genes were subcloned using 20 cycles of PCR to reduce the frequency of PCR artefacts.

In vitro mutagenesis

- In vitro mutagenesis was performed by PCR using recombinant pMAL-c with $Bet\ v\ 1$ inserted as template. Each mutant $Bet\ v\ 1$ gene was generated by 3 PCR reactions using 4 primers.
- mutation-specific oligonucleotide primers 15 synthesised accommodating each mutation, one for each DNA see Figs. 1 and 2. Usinq the mutated strand, nucleotide(s) as starting point both primers were extended 7 nucleotides in the 5'-end and 15 nucleotides in the 3'-end. The extending nucleotides were identical 20 in sequence to the $Bet\ v\ 1$ gene in the actual region.

Two generally applicable primers (denoted "all-sense" and "all non-sense" in Figure 2) were furthermore synthesised and used for all mutants. These primers were 15 nucleotides in length and correspond in sequence to regions of the pMAL-c vector approximately 1 kilobase upstream and downstream from the $Bet\ v\ 1$. The sequence of the upstream primer is derived from the sense strand and the sequence of the downstream primer is derived from the non-sense strand, see Fig. 2.

Two independent PCR reactions were performed essentially according to standard procedures (Saiki et al 1988, ref. 20) with the exception that only 20 temperature cycles were performed in order to reduce the frequency of PCR

artefacts. Each PCR reaction used pMAL-c with $Bet\ v\ 1$ inserted as template and one mutation-specific and one generally applicable primer in meaningful combinations.

Introduction of the four amino acid substitutions (Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly) in the Triple-patch mutant were performed like described above in a step by step process. First the Glu45Ser mutation then the Pro108Gly mutation and last the Asn28Thr, Lys32Gln mutations were introduced using pMAL-c with inserted Bet v 1 No. 2801, Bet v 1 (Glu45Ser), Bet v 1 (Glu45Ser, Pro108Gly) as templates, respectively.

The PCR products were purified by agarose gel electrophoresis and electro-elution followed by ethanol precipitation. A third PCR reaction was performed using the combined PCR products from the first two PCR reactions as template and both generally applicable primers. Again, 20 cycles of standard PCR were used. The PCR product was purified by agarose gel electrophoresis and electro-elution followed by ethanol precipitation, cut with restriction enzymes (BsiWI/EcoRI), and ligated directionally into pMAL-c with Bet v 1 inserted restricted with the same enzymes.

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Figure 3 shows an overview of all 9 $Bet\ v$ 1 mutations, which are as follows

Thr10Pro, Asp25Gly, Asn28Thr + Lys32Gln, Glu45Ser,

Asn47Ser, Lys55Asn, Glu60Ser (non-patch), Thr77Ala and
Pro108Gly. An additional mutant with four mutations was
also prepared (Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly).

Of these, five mutants were selected for further testing:
Asn28Thr + Lys32Gln, Glu45Ser, Glu60Ser, Pro108Gly and
the Triple-patch mutant Asn28Thr, Lys32Gln, Glu45Ser,
Pro108Gly.

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Nucleotide sequencing

Determination of the nucleotide sequence of the $Bet\ v\ 1$ encoding gene was performed before and after subcloning, and following in vitro mutagenesis, respectively.

Plasmid DNA's from 10 ml of bacterial culture grown to saturation overnight in LB medium supplemented with 0.1 g/l ampicillin were purified on Qiagen-tip 20 columns and sequenced using the Sequenase version 2.0 DNA sequencing kit (USB) following the recommendations of the suppliers.

Expression and purification of recombinant Bet v 1 and mutants

Recombinant Bet v 1 (Bet v 1 No. 2801 and mutants) were over-expressed in Escherichia coli DH 5a fused to maltose-binding protein and purified as described in ref. 15. Briefly, recombinant *E.coli* cells were grown at 37°C to an optical density of 1.0 at 436 nm, whereupon expression of the $Bet\ v\ 1$ fusion protein was induced by addition of IPTG. Cells were harvested by centrifugation 3 hours post-induction, re-suspended in lysis buffer and broken by sonication. After sonication and additional centrifugation, recombinant fusion protein was isolated amylose affinity chromatography and subsequently cleaved by incubation with Factor Xa (ref. 15). After F cleavage, recombinant Bet v 1 was isolated by gelfiltration and if found necessary, subjected to another round of amylose affinity chromatography in order to remove trace amounts of maltose-binding protein.

Purified recombinant Bet v 1 was concentrated by ultrafiltration to about 5 mg/ml and stored at 4 °C. The final yields of the purified recombinant Bet v 1

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preparations were between 2-5 mg per litre *E. coli* cell culture.

The purified recombinant Bet v 1 preparations appeared as single bands after silver-stained SDS-polyacrylamide electrophoresis with an apparent molecular weight of 17.5 kDa. N-terminal sequencing showed the expected sequences as derived from the cDNA nucleotide sequences and quantitative amino acid analysis showed the expected amino acid compositions.

We have previously shown (ref. 15) that recombinant $Bet\ v$ 1 No. 2801 is immunochemically indistinguishable from naturally occurring $Bet\ v$ 1.

Immunoelectrophoresis using rabbit polyclonal antibodies

The seven mutant $Bet\ v\ 1$ were produced as recombinant $Bet\ v\ 1$ proteins and purified as described above and tested for their reactivity towards polyclonal rabbit antibodies raised against $Bet\ v\ 1$ isolated from birch pollen. When analysed by immunoelectrophoresis (rocket-line immunoelectrophoresis) under native conditions, the rabbit antibodies were able to precipitate all mutants, indicating that the mutants had conserved α -carbon backbone tertiary structure.

In order to analyse the effect on human polyclonal IgE-response, the mutants Glu45Ser, Pro108Gly, Asn28Thr+Lys32Gln and Glu60Ser were selected for further analysis.

Bet v 1 Glu45Ser mutant

35 Glutamic acid in position 45 show a high degree of solvent-exposure (40%) and is located in a molecular

surface patch common for Fagales allergens (patch I). A serine residue was found to occupy position 45 in some of the Bet v 1 homologous PR-10 proteins arguing for that glutamic acid can be replaced by serine without distortion of the α -carbon backbone tertiary structure. In addition, as none of the known Fagales allergen sequences have serine in position 45, the substitution of glutamic acid with serine gives rise to a non-naturally occurring Bet v 1 molecule.

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T cell proliferation assay using recombinant Glu45Ser Bet v 1 mutant

The analysis was carried out as described in Spangfort et al 1996a. It was found that recombinant $Bet\ v\ 1$ Glu45Ser mutant was able to induce proliferation in T cell lines from three different birch pollen allergic patients with stimulation indices similar to recombinant and naturally occurring.

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Crystallisation and structural determination of recombinant Glu45Ser Bet v 1

Crystals of recombinant Glu45Ser Bet v 1 were grown by vapour diffusion at 25°C, essentially as described in (Spangfort et al 1996b, ref. 21). Glu45Ser Bet v 1, at a concentration of 5 mg/ml, was mixed with an equal volume of 2.0 M ammonium sulphate, 0.1 M sodium citrate, 1% (v/v) dioxane, pH 6.0 and equilibrated against 100x volume of 2.0 M ammonium sulfate, 0.1 M sodium citrate, After (v/v)dioxane, Нq 6.0. 24 hours equilibration, crystal growth was induced by applying the seeding technique described in ref. 21, using crystals of recombinant wild-type Bet v 1 as a source of seeds.

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After about 2 months, crystals were harvested and

analysed using X-rays generated from a Rigaku rotating anode as described in ref. 21 and the structure was solved using molecular replacement.

5 Structure of Bet v 1 Glu45Ser mutant

The structural effect of the mutation was addressed by growing three-dimensional $Bet\ v\ 1$ Glu45Ser protein crystals diffracting to 3.0 Å resolution when analysed by X-rays generated from a rotating anode. The substitution of glutamic acid to serine in position 45 was verified by the $Bet\ v\ 1$ Glu45Ser structure electron density map which also showed that the overall α -carbon backbone tertiary structure is preserved.

IgE-binding properties of Bet v 1 Glu45Ser mutant

The IgE-binding properties of $Bet\ v\ 1$ Glu45Ser mutant was compared with recombinant $Bet\ v\ 1$ in a fluid-phase IgE-inhibition assay using a pool of serum IgE derived from birch allergic patients.

Recombinant Bet v 1 no. 2801 was biotinylated at a molar ratio of 1:5 (Bet v 1 no. 2801:biotin). The inhibition assay was performed as follows: a serum sample (25 μ l) was incubated with solid phase anti IgE, washed, resuspended and further incubated with a mixture of biotinylated Bet v 1 no. 2801 (3.4 nM) and a given mutant (0-28.6 nM). The amount of biotinylated Bet v 1 no. 2801 bound to the solid phase was estimated from the measured RLU after incubation with acridinium ester labelled streptavidin. The degree of inhibition was calculated as the ratio between the RLU's obtained using buffer and mutant as inhibitor.

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Figure 4 shows the inhibition of the binding of

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biotinylated recombinant $Bet\ v\ 1$ to serum IgE from a pool of allergic patients by non-biotinylated $Bet\ v\ 1$ and by $Bet\ v\ 1$ Glu45Ser mutant.

There is a clear difference in the amount of respective 5 recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Bet v 1 reaches 50% inhibition at about 6.5 ng whereas the corresponding concentration for Bet v 1 10 Glu45Ser mutant is about 12 ng. This show that the point mutation introduced in Bet v 1 Glu45Ser mutant lowers the affinity for specific serum IgE by a factor of about 2. The maximum level of inhibition reached by the $\textit{Bet}\ \textit{v}\ \textit{1}$ Glu45Ser mutant is clearly lower compared to recombinant Bet v 1. This may indicate that after the Glu45Ser 15 substitution, some of the specific IqE present in the serum pool are unable to recognise the Bet v 1 Glu45Ser mutant.

20 Bet v 1 mutant Asn28Thr+Lys32Gln

Aspartate and lysine in positions 28 and 32, respectively show a high degree of solvent-exposure (35% and 50%, respectively) and are located in a molecular surface patch common for Fagales allergens (patch II). In the structure, aspartate 28 and lysine 32 are located close to each other on the molecular surface and most likely interact via hydrogen bonds. A threonine and a gluatamate residue were found to occupy positions 28 and 32, respectively in some of the Bet v 1 homologous PR-10 proteins arguing for that aspartate and lysine can be replaced with threonine and glutamate, respectively without distortion of the α -carbon backbone tertiary addition, structure. In as none of the naturally occurring isoallergen sequences have threonine and glutamate in positions 28 and 32, respectively, the

substitutions gives rise to a non-naturally occurring Bet v 1 molecule.

IgE-binding properties of Bet v 1 mutant

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The IgE-binding properties of mutant Asn28Thr+Lys32Gln was compared with recombinant $Bet\ v\ 1$ in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 5 shows the inhibition of the binding of biotinylated recombinant $Bet\ v\ 1$ to serum IgE from a pool of allergic patients by non-biotinylated $Bet\ v\ 1$ and by $Bet\ v\ 1$ mutant Asn28Thr+Lys32Gln.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Bet v 1 reaches 50% inhibition at about 6.5 ng whereas the corresponding concentration for Bet v 1 mutant Asn28Thr+Lys32Gln is about 12 ng. This show that the point mutations introduced in Bet v 1 mutant Asn28Thr+Lys32Gln lowers the affinity for specific serum IgE by a factor of about 2.

The maximum level of inhibition reached by the Bet v 1 mutant Asn28Thr+Lys32Gln mutant is clearly lower compared to recombinant Bet v 1. This may indicate that after the Asn28Thr+Lys32Gln substitutions, some of the specific IgE present in the serum pool are unable to recognise the Bet v 1 mutant Asn28Thr+Lys32Gln.

Bet v 1 mutant Pro108Gly

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Proline in position 108 show a high degree of solvent-

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exposure (60%) and is located in a molecular surface patch common for Fagales allergens (patch III). A glycine residue was found to occupy position 108 in some of the Bet v 1 homologous PR-10 proteins arguing for that proline can be replaced with glycine without distortion of the α -carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have glycine in position 108, the substitution of proline with glycine gives rise to a non-naturally occurring Bet v 1 molecule.

IgE-binding properties of Bet v 1 Pro108Gly mutant

The IgE-binding properties of $Bet\ v\ 1$ Pro108Gly mutant was compared with recombinant $Bet\ v\ 1$ in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 6 shows the inhibition of the binding of biotinylated recombinant $Bet\ v\ 1$ to serum IgE from a pool of allergic patients by non-biotinylated $Bet\ v\ 1$ and by $Bet\ v\ 1$ Pro108Gly mutant.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant $Bet\ v\ 1$ reaches 50% inhibition at about 6.5 ng whereas the corresponding concentration for $Bet\ v\ 1$ Pro108Gly is 15 ng. This show that the single point mutation introduced in $Bet\ v\ 1$ Pro108Gly lowers the affinity for specific serum IgE by a factor of about 2.

The maximum level of inhibition reached by the $Bet\ v\ 1$ Prologgly mutant is somewhat lower compared to recombinant $Bet\ v\ 1$. This may indicate that after the Prologgly substitution, some of the specific IqE present

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in the serum pool are unable to recognise the $Bet\ v\ 1$ PrologGly mutant.

Bet v 1 mutant Glu60Ser (non-patch mutant)

Glutamic acid in position 60 show a high degree of solvent-exposure (60%) however, it is not located in a molecular surface patch common for Fagales allergens. A serine residue was found to occupy position 60 in some of the Bet v 1 homologous PR-10 proteins arguing for that glutamic acid can be replaced with serine without distortion of the α -carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have serine in position 60, the substitution of glutamic acid with serine gives rise to a non-naturally occurring Bet v 1 molecule.

IgE-binding properties of Bet v 1 Glu60Ser mutant

- The IgE-binding properties of $Bet\ v\ 1$ Glu60Ser mutant was compared with recombinant $Bet\ v\ 1$ in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.
- 25 Figure shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Glu60Ser mutant. In contrast to the Glu45Ser, Pro108Gly and Asn28Thr+Lys32Gln mutants, the substitution glutamic acid 30 60 to serine, does not shown significant effect on the IgE-binding properties of. This indicates that substitutions outside the defined Fagales common patches only have a marginal effect on the binding specific serum IgE supporting the concept 35 conserved allergen molecular surface areas dominant IgE-binding epitopes.

Bet v 1 Triple-patch mutant

In the Triple-patch mutant, the point mutations

(Glu45Ser, Asn28Thr+Lys32Gln and Pro108Gly) introduced in the three different common Fagales patches, described above, were simultaneously introduced in creating an artificial mutant carrying four amino acid substitutions.

10 Structural analysis of Bet v 1 Triple-patch mutant

The structural integrity of the purified Triple-patch mutant was analysed by circular dichroism (CD) spectroscopy. Figure 8 shows the CD spectra recombinant and Triple-patch mutant, recorded at close to equal concentrations. The overlap in peak amplitudes and positions in the CD spectra from the two recombinant proteins shows that the two preparations contain equal amounts of secondary structures strongly suggesting that the α -carbon backbone tertiary structure is not affected by the introduced amino acid substitutions.

IgE-binding properties of $Bet\ v\ 1$ Triple-patch mutant

- The IgE-binding properties of $Bet\ v\ 1$ Triple-patch mutant was compared with recombinant $Bet\ v\ 1$ in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.
- 30 Figure 9 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Triple-patch mutant. In contrast to the single mutants described above, the inhibition curve of the 35 Triple-patch mutant is no longer parallel relative to

recombinant. This shows that the substitutions introduced

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in the Triple-patch mutant has changed the IgE-binding properties and epitope profile compared to recombinant. The lack of parallellity makes it difficult to quantify the decrease of the Triple-patch mutant affinity for specific serum IgE.

Recombinant Bet v 1 reaches 50% inhibition at about 6 ng whereas the corresponding concentration for Bet v 1 Triple-patch mutant is 30 ng, i.e a decrease in affinity by a factor 5. However, in order to reach 80% inhibition the corresponding values are 20 ng and 400 ng, respectively, i.e a decrease by a factor 20.

T cell proliferation assay using recombinant Bet v 1 Triple-patch mutant

The analysis was carried out as described in ref. 15. It was found that recombinant $Bet\ v\ 1$ Triple-patch mutant was able to induce proliferation in T cell lines from three different birch pollen allergic patients with stimulation indices similar to recombinant and naturally occurring. This suggests that the Triple-patch mutant can initiate the cellular immune response necessary for antibody production.

EXAMPLE 2

Example 2 describes the preparation of recombinant mutant allergens with one primary mutation. Recombinant mutant allergens according to the invention, i.e. allergens comprising at least four primary mutations, may be prepared using the same procedures.

Identification of common epitopes within *Vespula vulgaris*venom major allergen antigen <u>5</u>

Antigen 5 is one of the three vespid venom proteins, which are known allergens in man. The vespids include hornets, yellow-jacket and wasps. The other two known allergens of vespid venoms are phospholipase A1 and hyaluronidase. Antigen 5 from Vespula vulgaris (Ves v 5) has been cloned and expressed as recombinant protein in the yeast system (Monsalve et al. 1999, ref. 22). The three-dimensional crystal structure of recombinant Ves v 5 has recently been determined at 1.8 Å resolution (in preparation). The main features of the structure consist of four $\beta\text{-strands}$ and four $\alpha\text{-helices}$ arranged in three stacked layers giving rise to a " α - β - α sandwich". The sequence identity between Antigen 5 homologous allergens from different Vespula species is about 90% suggesting presence of conserved molecular surface areas and B cell epitopes.

The presence and identification of common patches was performed after alignment of all known amino acid sequences, previously described as for tree pollen allergens, of the *Vespula* antigen 5 allergens combination with an analysis of the molecular surface of Antigen 5 revealed by the three-dimensional structure of Ves v 5. Figure 10 shows solvent accessibility of individually aligned antigen 5 residues and alignment of Vespula antigen 5 sequences (left panel). On the right panel of figure 10 is shown the molecular surface of antigen 5 with conserved areas among Vespula antigen 5:s coloured.

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Selection of amino acid residues for site-directed mutagenesis

Amino acid residues for site-directed mutagenesis were selected among residues present the patches common for Vespula since modifications of these is expected to

affect the binding of serum IgE from the majority of patients showing clinical *Vespula* allergic cross-reactivity.

The relative orientation and percentage of solvent-exposure of each amino acid residue within respective patch was calculated based on their atomic coordinates. Residues having a low degree of solvent exposure were not regarded suitable for mutagenesis due to the possible disruption of the structure or lack of antibody interaction. The remaining residues were ranked according to their degree of solvent-exposure.

Cloning of the gene encoding Ves v = 5

Total RNA was isolated from venom acid glands of *Vespula vulgaris* vespids as described in (Fang *et al.* 1988, ref. 23).

20 First-strand cDNA synthesis, PCR amplification and cloning of the Ves v 5 gene was performed as described in (Lu et al. 1993, ref. 24)

Subcloning into pPICZαA

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The gene encoding $Ves\ v$ 5 was subsequently sub-cloned into the pPICZ α A vector (Invitrogen) for secreted expression of $Ves\ v$ 5 in $Pichia\ pastoris$. The gene was amplified by PCR and sub-cloned in frame with the coding sequence for the α -factor secretion signal of $Saccharomyces\ cerevisiae$. In this construct the α -factor is cleaved off, in vivo, by the $Pichia\ pastoris\ Kex2$ protease system during secretion of the protein.

In brief PCR was performed using $Ves\ v\ 5$ as template and primers corresponding to the amino- and carboxyterminus

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of the protein, respectively. The primers were extended in the 5'-end to accommodate restriction sites for cloning, EcoRI and XbaI, respectively. Nucleotides encoding the Kex2 cleavage site was in this construct positioned 18 nucleotides upstream to the amino terminus of the protein, resulting in the expression of Ves V 5 with six additional amino acids, Glu-Ala-Glu-Ala-Glu-Phe, at the amino terminus.

10 Insertion of pPICZαA-Ves v 5 into P. pastoris

The pPICZ α A vectors with the *Ves v 5* gene inserted was linearised by Sac I restriction and inserted into the *AOX1* locus on the *Pichia pastoris* genome. Insertion was performed by homologous recombination on *Pichia pastoris* KM71 cells following the recommendations of Invitrogen.

In vitro mutagenesis

- 20 In vitro mutagenesis was performed by PCR using recombinant pPICZ α A with Ves v 5 inserted as template. Each mutant Ves v 5 gene was generated by 3 PCR reactions using 4 primers.
- 25 mutation-specific oligonucleotide primers synthesised accommodating each mutation, one for each DNA see Figures 11 and 12. Using the mutated strand, both primers nucleotide(s) as starting point were in 5'-end and 12-13 extended 6-7 nucleotides the nucleotides in the 3'-end. The extending nucleotides were 30 identical in sequence to the $Ves \ v \ 5$ gene in the actual region.

Two generally applicable primers (denoted "all sense" and "all non-sense" in Figure 12) were furthermore synthesised and used for all mutants. To insure

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expression of Ves v 5 mutants with authentic amino terminus, one primer corresponding to the amino terminus of the protein was extended in the 5'-end with a Xho I site. Upon insertion of the Ves v 5 mutant genes into the pPICZαA vector, the Kex2 protease cleavage site was regenerated directly upstream to the amino terminus of Ves v 5. The second primer was corresponding in sequence region of the pPICZαA vector positioned approximately 300 bp downstream from the Ves v 5 gene. The sequence of the primer corresponding to the amino terminus of Ves v 5 is derived from the sense strand and the sequence of the downstream primer is derived from the non-sense strand, see Figure 11.

Two independent PCR reactions were performed essentially according to standard procedures (Saiki et al 1988) with the exception that only 20 temperature cycles were performed in order to reduce the frequency of PCR artefacts. Each PCR reaction used pPICZαA with Ves v 5 inserted as template and one mutation-specific and one generally applicable primer in meaningful combinations.

The PCR products were purified by using "Concert, Rapid PCR Purification System" (Life Technologies). A third PCR reaction was performed using the combined PCR products from the first two PCR reactions as template and both generally applicable primers. Again, 20 cycles of standard PCR were used. The PCR product was purified with the "Concert, Rapid PCR Purification System" (Life Technologies), cut with restriction enzymes (XhoI/XbaI), and ligated directionally into pPICZ α A vector restricted with the same enzymes. Figure 13 shows an overview of all Ves v 5 mutations.

Insertion of pPICZ α A-Ves v 5 mutants into P. pastoris

The pPICZ α A vectors with the *Ves v 5 mutant* genes inserted were linearised by Sac I restriction and inserted into the *AOX1* locus on the *Pichia pastoris* genome. Insertions were performed by homologous recombination on *Pichia pastoris* KM71 cells following the recommendations of Invitrogen.

Nucleotide sequencing

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Determination of the nucleotide sequence of the Ves v 5 encoding gene was performed before and after subcloning, and following in vitro mutagenesis, respectively.

Plasmid DNA's from 10 ml of bacterial culture grown to saturation overnight in LB medium supplemented with 0.1 g/l ampicillin were purified on Qiagen-tip 20 columns and sequenced using the Sequenase version 2.0 DNA sequencing kit (USB) following the recommendations of the suppliers.

20 Expression and purification of recombinant $Ves \ v$ 5

Recombinant yeast cells of *Pichia pastoris* strain KM71 were grown in 500 ml bottles containing 100 ml of pH 6.0 phosphate buffer containing yeast nitrogen base, biotin, glycerol and histidine at 30° C with orbital shaking at 225 rpm until A_{600} nm of 4-6. Cells were collected by centrifugation and re-suspended in 10 ml of similar buffered medium containing methanol in place of glycerol. Incubation was continued at 30° C for 7 days with daily addition of 0.05 ml methanol.

Cells were harvested by centrifugation and the collected culture fluid was concentrated by ultrafiltration. After dialysis against 50 mM ammonium acetate buffer, pH 4.6, the sample was applied to a FPLC (Pharmacia) SE-53 cation exchange column equilibrated in the same buffer. The

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column was eluated with a 0-1.0 M NaCl, 50 mM ammonium acetate linear gradient. The recombinant Ves v 5 peak eluting at about 0.4 M NaCl was collected and dialysed against 0.02 N acetic acid. After concentration to about 10 mg/ml, the purified Ves v 5 was stored at 4° C.

Crystallisation of recombinant Ves v 5

Crystals of Ves v 5 was grown by the vapour diffusion technique at 25°C. For crystallisation, 5 μ l of 5 mg/ml Ves v 5 was mixed with 5 μ l of 18% PEG 6000, 0.1 M sodium citrate, pH 6.0 and equilibrated against 1 ml of 18% PEG 6000, 0.1 M sodium citrate, pH 6.0.

15 X-ray diffraction data was collected at 100K from native Ves v 5 crystals and after incorporation of heavy-atom derivatives and used to solve the three-dimensional structure of Ves v 5, see Figure 10 (manuscript in preparation).

Immunoelectrophoresis using rabbit polyclonal antibodies

The two $Ves\ v$ 5 mutants were produced as recombinant $Ves\ v$ 5 proteins and tested for their reactivity towards polyclonal rabbit antibodies raised against recombinant $Ves\ v$ 5. When analysed by rocket immunoelectrophoresis under native conditions, the rabbit antibodies were able to precipitate recombinant $Ves\ v$ 5 as well as both mutants, indicating that the mutants have conserved α -carbon backbone tertiary structure.

Inhibition of specific serum IqE

The IgE-binding properties of $Ves\ v\ 5$ mutants were compared to recombinant $Ves\ v\ 5$ in a fluid-phase IgE-inhibition assay using a pool of serum IgE derived from

vespid venom allergic patients.

The inhibition assay was performed as described above using biotinylated recombinant $Ves\ v$ 5 instead of $Bet\ v$ 1.

Ves v 5 Lys72Ala mutant

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Lysine in position 72 show a high degree of solventexposure (70%) and is located in a molecular surface patch common for Vespula antigen 5. The relative orientation and high degree of solvent exposure argued for that lysine 72 can be replaced by an alanine residue without distortion of the α -carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have alanine in position 72, the substitution of lysine with alanine gives rise to a non-naturally occurring $Ves\ v$ 5 molecule.

20 IgE-binding properties of Ves v 5 Lys72Ala mutant

The IgE-binding properties of $Ves\ v$ 5 Lys72Ala mutant was compared with recombinant $Ves\ v$ 5 in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 14 shows the inhibition of the binding of biotinylated recombinant $Ves\ v\ 5$ to serum IgE from a pool of allergic patients by non-biotinylated $Ves\ v\ 5$ and by $Ves\ v\ 5$ Lys72Ala mutant.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant $Ves\ v\ 5$ reaches 50% inhibition at about 6 ng whereas the corresponding concentration for $Ves\ v\ 5$

Lys72Ala mutant is 40 ng. This show that the single point mutation introduced in Ves v 5 Lys72Ala mutant lowers the affinity for specific serum IgE by a factor of about 6. The maximum level of inhibition reached by the Ves v 5 Lys72Ala mutant significantly lower compared to recombinant Ves v 5. This may indicate that after the Lys72Ala substitution, some of the specific IgE present in the serum pool are unable to recognise the Ves v 5 Lys72Ala mutant.

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Ves v 5 Tyr96Ala mutant

Tyrosine in position 96 show a high degree of solvent-exposure (65%) and is located in a molecular surface patch common for *Vespula* antigen 5. The relative orientation an high degree of solvent exposure argued for that tyrosine 96 can be replaced by an alanine residue without distortion of the three-dimensional structure. In addition, as none of the naturally occurring isoallergen sequences have alanine in position 96, the substitution of tyrosine with alanine gives rise to a non-naturally occurring *Ves v 5* molecule.

IgE-binding properties of $Ves\ v$ 5 Tyr96Ala mutant

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The IgE-binding properties of $Ves\ v\ 5$ Tyr96Ala mutant was compared with recombinant $Ves\ v\ 5$ in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

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Figure 14 shows the inhibition of the binding of biotinylated recombinant $Ves\ v$ 5 to serum IgE from a pool of allergic patients by non-biotinylated $Ves\ v$ 5 and by $Ves\ v$ 5 Tyr96Ala mutant.

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There is a clear difference in the amount of respective

recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant $Ves\ v$ 5 reaches 50% inhibition at about 6 ng whereas the corresponding concentration for $Ves\ v$ 5 Tyr96Ala mutant is 40 ng.

This show that the single point mutation introduced in $Ves\ v$ 5 Tyr96Ala mutant lowers the affinity for specific serum IgE by a factor of about 6.

The maximum level of inhibition reached by the $Ves\ v\ 5$ Tyr96Ala mutant significantly lower compared to recombinant $Ves\ v\ 5$. This may indicate that after the Tyr96Ala substitution, some of the specific IgE present in the serum pool are unable to recognise the $Ves\ v\ 5$ Tyr96Ala mutant.

EXAMPLE 3

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20 <u>Identification and selection of amino acids for</u> substitution

The parameters of solvent accessibility and conservation degree were used to identify and select surface-exposed amino acids suitable for substitution for the allergens Bet v 1, Der p 2 and Ves v 5.

Solvent accessibility

30 Solvent accessibility was calculated using the software InsightII, version 97.0 (MSI) and a probe radius of 1.4 Å (Connolly surface).

Internal cavities were excluded from the analyses by filling with probes using the software PASS (Putative Active Sites with Spheres). Probes on the surface were subsequently removed manually.

Conservation

5 <u>Bet v 1:</u>

3-D structure is based on accession number Z80104 (1bvl.pdb).

10 38 other Bet v 1 sequences included in the analysis of conserved residues comprise accession numbers: P15494=X15877=Z80106, Z80101, AJ002107, Z72429, AJ002108, Z80100, Z80103, AJ001555, Z80102, Z80105, Z72436, P43183=X77271, Z72430, AJ002106, P43178=X77267, P43179=X77268, P43177=X77266, Z72438, 15 P43180=X77269, AJ001551, P43185=X77273, AJ001557, Z72434, Z72433=P43186, AJ001554, X81972, Z72431, P45431=X77200, P43184=X77272, P43176=X77265, S47250, S47251, Z72435,

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Der p 2:

- 3-D structure is based on accession number P49278 (la9v.pdb).
- 6 other Der p 2 sequences included in the analysis of conserved residues comprise the following substitutions: ALK-G: V40L, T47S, M111L, D114N.

ALK-101: M76V.

ALK-102: V40L, T47S.

30 ALK-104: T47S, M111I, D114N.

Z72439, Z72437, S47249.

ALK-113: T47S.

ALK-120: V40L, T47S, D114N.

Ves v 5:

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3-D structure is based on accession number Q05110 (pdb

coordinates unpublished).

Another Ves v 5 sequence in the analysis of conserved residues contains one amino acid substitution: M202K.

Results

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Bet v 1

10 59 amino acids highly solvent exposed:
 K-129, E-60, N-47, K-65, P-108, N-159, D-93, K-123, K-32,
 D-125, R-145, D-109, T-77, E-127, Q-36, E-131, L-152, E-6, E-96, D-156, P-63, H-76, E-8, K-134, E-45, T-10, V-12,
 K-20, L-62, S-155, H-126, P-50, N-78, K-119, V-2, L-24,
15 E-42, N-4, A-153, I-44, E-138, G-61, A-130, R-70, N-28,
 P-35, S-149, K-103, Y-150, H-154, N-43, A-106, K-115, P-14, Y-5, K-137, E-141, E-87, E-73.

57 amino acids highly solvent exposed and conserved (>70%):

K-129, E-60, N-47, K-65, P-108, N-159, D-93, K-123, K-32, D-125, R-145, D-109, E-127, Q-36, E-131, L-152, E-6, E-96, D-156, P-63, H-76, E-8, K-134, E-45, T-10, V-12, K-20, S-155, H-126, P-50, N-78, K-119, V-2, L-24, E-42, N-4, A-153, I-44, E-138, G-61, A-130, R-70, N-28, P-35, S-149, K-103, Y-150, H-154, N-43, A-106, K-115, P-14, Y-5, K-137, E-141, E-87, E-73.

23 mutations performed:

30 Y5V, T10P, D25E, N28T, K32Q, E42S, E45S, N47S, K55N, K65N, T77A, N78K, E96L, K97S, K103V, P108G, D109N, K123I, D125Y, K134E, R145E, D156H, +160N.

Table 1 shows a listing in descending order of solvent exposure of Bet v 1 amino acids. Column 1 lists the amino acid number starting from the amino-terminal, column 2

lists the amino acid in one letter abbreviation, column 3 lists the normalised solvent exposure index, column 4 lists the percent of known sequences having the concerned amino acid in this position.

Table 1: Bet v 1

5

NO	AA	Solv_exp Cons	%
	129K	1,000	90
	60E	0,986	97
	47N	0,979	100
	65K	0,978	100
	108P	0,929	100
	159N	0,869	100
	93D	0,866	100
	123K	0,855	100
	32K	0,855	100
	125D	0,821	74
	145R	0,801	90
	109D	0,778	82
	7 7 T	0,775	56
	127E	0,760	100
	36Q	0,749	95
	131E	0,725	100
	152L	0,718	97
	6E	0,712	100
	96E	0,696	100
	156D	0,693	97
	63P	0,692	97
	76H	0,683	90
	8E	0,638	97
	134K	0,630	100
	45E	0,623	100
	10T	0,613	97
	12V	0,592	100
	20K	0,584	100
	62L	0,575	5
	155S	0,568	97
	126H	0,551	95
	50P	0,541	100
	78N	0,538	100
	119K	0,529	100
	2V	0,528	100
	24L	0,528	100
	42E	0,519	100
	4N	0,517	95
	153A	0,513	100
	441	0,508	97
	138E	0,496	100
	61G	0,488	100

130A	0,479	97
70R	0,474	100
28N	0,469	90
35P	0,467	100
149S	0,455	92
103K	0,447	100
150Y	0,438	100
154H	0,436	100
43N	0,412	100
106A	0,411	95
115K 14P	0,411 0.410	100 97
5Y	0,410	100
137K	0,396	100
141E	0,387	95
87E	0,385	100
73E	0,384	100
16A	0,367	100
79F	0,362	100
3F	0,355	100
158Y	0,346	100
105V	0,336	100
101E	0,326	100
64F	0,325	100
861	0,322	100
39S	0,314	100
124G	0,310	100
72D	0,308	97
142T	0,293	67 100
66Y 55K	0,289 0,288	100 100
7T	0,288	67
40S	0,274	95
25D	0,271	87
135A	0,267	92
68K	0,262	100
97K	0,247	100
46G	0,235	100
27D	0,232	97
1G	0,227	100
1131	0,225	7 7
51G	0,220	100
92G	0,218	100
80K	0,212	100
110G	0,211	100
107T	0,203	85 92
94T 41V	0,202 0,201	92 97
48G	0,198	100
911	0,198	18
31P	0,188	100
75D	0,188	97
33V	0,183	100
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49G	0,176	100
17R	0,172	100
99S	0,158	64
89G	0,154	100
531	0,154	100
121H	0,153	100
9T	0,150	72
74V	0,148	97
132Q	0,146	72
57S	0,137	49
148E	0,135	100
82N	0,133	41
128V	0,125	64
117S	0,124	87
90P	0,117	67
1161	0,112	100
122T	0,107	100
139M	0,104	62
95L	0,104	97
54K	0,096	100
146A	0,095	100
59P	0,088	97
157A	0,088	100
133V	0,077	44
88G	0,068	100
140G	0,053	85
37A	0,042	95
81Y	0,041	100
231	0,036	95
1041	0,036	92
15A	0,036	97
58F	0,029	100
29L	0,028	100
19F	0,027	100
100N	0,022	97
22F	0,021	97
71V	0,014	100
111G	0,014	100
131	0,014	100
18L	0,014	97
114L	0,014	100
11S	0,007	100
151L	0,007	97
144L	0,007	90
52T	0,007	100
84S	0,007	97
118N	0,007	97
1021	0,007	100
21A	0,000	97
26G	0,000	97
30F 34A	0,000	44
	0,000	100
381	0,000	87

561	0,000	100
67V	0,000	97
69D	0,000	62
83Y	0,000	95
85V	0,000	72
981	0,000	95
1128	0,000	77
120Y	0,000	95
136S	0,000	67
143L	0,000	100
147V	0,000	100

Der p 2

55 amino acids highly solvent exposed:

5 R-128, D-129, H-11, H-30, S-1, K-77, Y-75, R-31, K-82, K-6, K-96, K-48, K-55, K-89, Q-85, W-92, I-97, H-22, V-65, S-24, H-74, K-126, L-61, P-26, N-93, D-64, I-28, K-14, K-100, E-62, I-127, E-102, E-25, P-66, D-114, L-17, G-60, P-95, E-53, V-81, K-51, N-103, Q-2, N-46, E-42, T-91, D-87, N-10, M-111, C-8, H-124, I-68, P-79, K-109, K-15.

54 amino acids highly solvent exposed and conserved (>70%):

R-128, D-129, H-11, H-30, S-1, K-77, Y-75, R-31, K-82, K-6, K-96, K-48, K-55, K-89, Q-85, W-92, I-97, H-22, V-65, S-24, H-74, K-126, L-61, P-26, N-93, D-64, I-28, K-14, K-100, E-62, I-127, E-102, E-25, P-66, L-17, G-60, P-95, E-53, V-81, K-51, N-103, Q-2, N-46, E-42, T-91, D-87, N-10, M-111, C-8, H-124, I-68, P-79, K-109, K-15.

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6 mutations performed:

K6A, K15E, H30N, E62S, H74N, K82N

Table 2 shows a listing in descending order of solvent
25 exposure of Der p 2 amino acids. Column 1 lists the amino
acid number starting from the amino-terminal, column 2
lists the amino acid in one letter abbreviation, column 3
lists the normalised solvent exposure index, column 4

lists the percent of known sequences having the concerned amino acid in this position.

Table 2: Der p 2

NO	AA	Solv_exp Cons	%
	128R	1,000	100
	129D	0,965	100
	11H	0,793	100
	30H	0,712	100
	18	0,700	100
	77K	0,694	
	75Y		100
	31R	0,681	100
		0,677	100
	82K 6K	0,658	100
		0,645	100
	96K	0,643	100
	48K	0,642	100
	55K	0,641	100
	89K	0,627	100
	85Q	0,624	100
	92W	0,610	100
	971	0,581	100
	22H	0,568	100
	65V	0,559	100
	24\$	0,557	100
	74H	0,542	100
	126K	0,542	100
	61L	0,539	100
	26P	0,516	100
	93N	0,513	100
	64D	0,509	100
	281	0,504	100
	14K	0,493	100
	100K	0,489	100
	62E	0,454	100
	1271	0,439	100
	102E	0,428	100
	25E	0,428	100
	66P	0,427	100
	114D	0,418	57
	17L	0,412	100
	60 G	0,390	100
	95P	0,388	100
	53E	0,377	100
	81V	0,377	100
	51K	0,370	100
	103N	0,369	100
	2Q	0,366	100
	46N	0,360	100
	42E	0,357	100
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32G	0,059	100
99P	0,058	100
86Y	0,056	100
16V	0,052	100
50A	0,040	100
90Y	0,039	100
18V	0,035	100
33K	0,033	100
521	0,029	100
581	0,029	100
104V	0,024	100
112G	0,023	100
21C	0,023	100
881	0,023	100
117L	0,016	100
56A	0,011	100
41F	0,011	100
120A	0,006	100
119C	0,006	100
67G	0,005	100
122A	0,005	100
37L	0,000	100
39A	0,000	100
43A	0,000	100
47T	0,000	29
80L	0,000	100
106V	0,000	100
108V	0,000	100
110V	0,000	100
118A	0,000	100
125A	0,000	100

Ves v 5

89 amino acids highly solvent exposed:

K-16, K-185, K-11, K-44, K-210, R-63, K-13, F-6, K-149, K-128, E-184, K-112, K-202, F-157, E-3, K-29, N-203, N-34, K-78, K-151, L-15, L-158, Y-102, W-186, K-134, D-87, K-52, T-67, T-125, K-150, Y-40, Q-48, L-65, K-81, Q-101, Q-208, K-144, N-8, N-70, H-104, Q-45, K-137, K-159, E-205, N-82, A-111, D-131, K-24, V-36, N-7, M-138, T-209, 10 V-84, K-172, V-19, D-56, P-73, G-33, T-106, N-170, L-28, T-43, Q-114, C-10, K-60, N-31, K-47, E-5, D-145, V-38, A-127, D-156, E-204, P-71, G-26, Y-129, D-141, F-201, R-68, N-200, D-49, S-153, K-35, S-39, Y-25, V-37, G-18, W-85,

15 I-182. 88 amino acids highly solvent exposed and conserved (>70%):

K-16, K-185, K-11, K-44, K-210, R-63, K-13, F-6, K-149, K-128, E-184, K-112, F-157, E-3, K-29, N-203, N-34, K-78, K-151, L-15, L-158, Y-102, W-186, K-134, D-87, K-52, T-67, T-125, K-150, Y-40, Q-48, L-65, K-81, Q-101, Q-208, K-144, N-8, N-70, H-104, Q-45, K-137, K-159, E-205, N-82, A-111, D-131, K-24, V-36, N-7, M-138, T-209, V-84, K-172, V-19, D-56, P-73, G-33, T-106, N-170, L-28, T-43, Q-114, C-10, K-60, N-31, K-47, E-5, D-145, V-38, A-127, D-156, E-204, P-71, G-26, Y-129, D-141, F-201, R-68, N-200, D-49, S-153, K-35, S-39, Y-25, V-37, G-18, W-85, I-182.

15 9 mutations performed:

K29A, T67A, K78A, V84S, Y102A, K112S, K144A, K202M, N203G

Table 3 shows a listing in descending order of solvent exposure of Ves v 5 amino acids. Column 1 lists the amino acid number starting from the amino-terminal, column 2 lists the amino acid in one letter abbreviation, column 3 lists the normalised solvent exposure index, column 4 lists the percent of known sequences having the concerned amino acid in this position.

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Table 3: Ves v 5

NO	AA	Solv_exp	
	16K	1,000	100
	185K	0,989	100
	11K	0,978	100
	44K	0,978	100
	210K	0,962	100
	63R	0,956	100
	13K	0,951	100
	6F	0,868	100
	149K	0,868	100
	128K	0,857	100
	184E	0,841	100
	112K	0,824	100

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202K	0,824	50
157F	0,819	100
3E	0,802	100
29K	0,797	100
203N	0,797	100
34N	0,775	100
78K	0,775	100
151K	0,753	100
15L	0,714	100
158L	0,714	100
102Y	0,687	100
186W	0,665	100
134K	0,654	100
87D	0,621	100
52K	0,615	100
67T	0,610	100
125T	0,610	100
150K	0,604	100
40Y	0,593	100
48Q	0,593	100
65L	0,593	100
81K	0,588	100
101Q	0,577	100
208Q	0,566	100
144K	0,560	100
8N 70N	0,555	100
70N	0,549	100
104H 45Q	0,549 0.538	100
137K	0,538 0,538	100 100
157K	0,533	100
205E	0,511	100
82N	0,500	100
111A	0,500	100
131D	0,495	100
24K	0,489	100
36V	0,489	100
7N	0,484	100
138M	0,473	100
209T	0,473	100
84V	0,462	100
172K	0,451	100
19V	0,445	100
56D	0,445	100
73P	0,440	100
33G	0,429	100
106T	0,429	100
170N	0,429	100
28L	0,423	100
43T	0,423	100
114Q	0,423	100
10C	0,412	100
60K	0,407	100

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31N	0,396	100
47K	0,396	100
5E	0,390	100
145D	0,390	100
38V	0,379	100
127A 156D	0,379	100
156D 204E	0,379	100
71P	0,374	100 100
26G	0,363 0,352	100
129Y	0,352	100
141D	0,341	100
201F	0,341	100
68R	0,335	100
200N	0,308	100
49D	0,302	100
153S	0,302	100
35K	0,297	100
39S	0,291	100
25Y	0,280	100
37V	0,280	100
18G	0,275	100
85W	0,275	100
1821	0,275	100
46E	0,264	100
126A	0,253	100
88E	0,247	100
76P 79N	0,236	100 100
79N 124S	0,236 0,236	100
30P	0,231	100
123G	0,231	100
162H	0,231	100
183Q	0,231	100
121	0,225	100
197P	0,225	100
130D	0,220	100
148P	0,214	100
180K	0,214	100
23C	0,209	100
75P	0,209	100
113Y	0,209	100
108R	0,203	100
188K	0,203	100
51L 59Q	0,198	100
121L	0,198 0,198	100 100
121E 122T	0,198	100
154G	0,192	100
53E	0,170	100
72G	0,170	100
41G	0,165	100
86N	0,165	100

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EXAMPLE 4

This Example describes preparation and characterisation of recombinant mutant Bet v 1 allergens according to the invention, i.e. allergens with diminished IgE-binding affinity comprising at least four primary mutations.

Selection of amino acid residues for site-directed mutagenesis of Bet v 1

Solvent accessibility of amino acid residues of Bet v 1 is shown in Example 3, table 1. The rate of amino acid conservation is based on sequence alignment performed at Molecular the ExPaSy Biology Server (http://www.expasy.ch/) using the ClustalW algorithm on a BLAST search where the Bet V 1.2801 wild type amino acid sequence is used as input sequence. The includes 67 allergen sequences (39 Bet v 1 sequences, 11 Car b 1 sequences, 6 Cor a 1 sequences, and 13 Aln g 1 sequences) from species within the order Fagales (Bet v 1: Betula verrrucosa; Car b 1: Carpinus betulus; Cor a 1: Corylus avellana; Aln g 1: alnus glutinosa). In respect to the mutated recombinant Bet v 1 allergens shown in the examples, target residues for substitution was based on ≥95% amino acid identity.

As described in Example 1, amino acid residues with a high degree of solvent-exposure and a high degree of conservation between pollen allergens from related species, were selected for site-directed mutagenesis. Residues having a low degree of solvent exposure (<20%) were not regarded relevant for mutagenesis due to the possible disruption of the tertiary structure or lack of antibody interaction.

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The introduced residues were all present in corresponding positions in isoforms of a group of plant proteins called related pathogenesis (PR-10) proteins. Molecular the tertiary structures of modelling suggests that Fagales allergens and PR-10 proteins are close to being identical. Bet v 1 shares significant sequence identity (20-40%) with PR-10 proteins. However, there are no reports of allergic cross-reactivity towards these PR-10 Thus, exchange of a highly conserved and solvent exposed amino acid from Bet v 1 with an amino acid in the corresponding position in a PR-10 protein,

results in a mutated Bet v 1 protein with an unaltered α -carbon backbone tertiary structure but with diminished IqE-binding affinity.

5 In vitro mutagenesis

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was performed by PCR vitro mutagenesis recombinant pMAL-c with Bet v 1 inserted as template. Preparation of recombinant mutant allergens comprising five to nine primary mutations included two PCR steps; step I and II. First, each single mutation (or several mutations if located closely together in the sequence) was introduced into sequential DNA sequences of Bet v 1.2801 or Bet v 1.2801 derivatives using sense and mutation-specific oligonucleotide anti-sense accommodating each mutation(s) along with sense and antioligonucleotide primers accommodating sense upstream or downstream neighbour mutations or the Nterminus/C-terminus of Bet ν 1, respectively schematically illustrated in Figure 17 (I). Secondly, PCR products from PCR reaction I were purified, mixed and used as templates for an additional PCR reaction (II) with oligonucleotide primers accommodating the N-terminus and C-terminus of Bet v 1 as schematically illustrated in Figure 17 (II). The PCR products were purified by agarose gel electrophoresis and PCR gel purification Techhnologies) followed by ethanol precipitation, with restriction enzymes (SacI/EcoRI) or (SacI/ XbaI), and ligated directionally into pMAL-c restricted with the same enzymes.

Figure 18 shows synthesised oligonucleotide primers and schematically illustrations for the construction of Bet v 1 mutants with five to nine primary mutations. The mutated amino acids were preferably selected from the group consisting of amino acids that are characterised by

being highly solvent exposed and conserved as described in Example 3. The Bet v 1 mutants are the following primary and secondary mutations stated in parenthesis:

5 Mutant Bet v 1 (2628): Tyr5Val, Glu45Ser, Lys65Asn, Lys97Ser, Lys134Glu.

Mutant Bet v 1 (2637): Ala16Pro, (Asn28Thr, Lys32Gln), Lys103Thr, Pro108Gly, (Leu152Lys, Ala153Gly, Ser155Pro).

Mutant Bet v 1 (2733): (Tyr5Val, Lys134Glu), (Asn28Thr, Lys32Gln), Glu45Ser, Lys65Asn, (Asn78Lys, Lys103Val), Lys97Ser, Pro108Gly, Arg145Glu, (Asp156His, +160Asn)

Mutant Bet v 1 (2744): (Tyr5Val, Lys134Glu), (Glu42Ser, Glu45Ser), (Asn78Lys, Lys103Val), Lys123Ile, (Asp156His, +160Asn).

Mutant Bet v 1 (2753): (Asn28Thr, Lys32Gln), Lys65Asn,

(Glu96Leu, Lys97Ser), (Pro108Gly, Asp109Asn), (Asp125Tyr,

Glu127Ser), Arg145Glu.

Nucleotide sequencing

Determination of the nucleotide sequence of the Bet v 1 encoding gene was performed before and after subcloning, and following in vitro mutagenesis, respectively.

Plasmid DNA's from 10 ml of bacterial culture grown to saturation overnight in LB medium supplemented with 0.1 g/l ampicillin were purified on Qiagen-tip 20 columns and sequenced using the Ready reaction dye terminator cycle sequencing kit and a Fluorescence Sequencer AB PRISM 377, both from (Perkin Elmer), following the recommendations of the suppliers.

Expression and purification of recombinant Bet v 1 and mutants

Recombinant Bet v 1 (Bet v 1.2801 and mutants) were overexpressed in Escherichia coli DH 5α fused to maltosebinding protein and purified as described in ref. 15. Briefly, recombinant E.coli cells were grown at 37°C to an optical density of 0.8 at 600 nm, whereupon expression of Bet v 1 fusion protein was induced by addition of IPTG. Cells were harvested by centrifugation 3 hours post-induction, re-suspended in lysis buffer and broken sonication. After sonication and additional centrifugation, recombinant fusion protein was isolated amylose affinity chromatography and subsequently cleaved by incubation with Factor Xa (ref. 15). After F Xa cleavage, recombinant Bet v 1 was isolated by gelfiltration and subjected to another round of amylose affinity chromatography in order to remove trace amounts of maltose-binding protein.

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Purified recombinant Bet v 1 was concentrated by ultrafiltration to about 5 mg/ml and stored at 4 $^{\circ}$ C. The final yields of the purified recombinant Bet v 1 preparations were between 2-5 mg per litre E. coli cell culture.

The purified recombinant Bet v 1 preparations appeared as single bands after silver-stained SDS-polyacrylamide electrophoresis with an apparent molecular weight of $17.5 \, \mathrm{kDa}$.

We have previously shown (ref. 15) that recombinant $Bet\ v$ 1 No. 2801 is immunochemically indistinguishable from naturally occurring Bet v 1.

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Figure 19 shows introduced point mutations at the molecular surface of Bet v 1 (2628) and Bet v 1 (2637). In mutant Bet v 1 (2628) five primary mutations were introduced in one half of Bet v 1 leaving the other half unaltered. In mutant Bet v 1(2637) five primary and three secondary mutations were introduced in the other half leaving the first half unaltered. In this way, mutations in mutant Bet v 1 (2628) and mutant Bet v 1(2637) affects different halves of the Bet v 1 surface, respectively.

Crystallisation and structural determination of recombinant Bet v 1(2628) mutant protein.

Crystals of recombinant Bet v 1 (2628) were grown by 15 vapour diffusion at 25°C, essentially as described in (Spangfort et al 1996b, ref. 21). Bet v 1 (2628), at a concentration of 5 mg/ml, was mixed with an equal volume of 2.2 M ammonium sulphate, 0.1 M sodium citrate, 1% (v/v) dioxane, pH 6.3 and equilibrated against 100x 20 volume of 2.2 M ammonium sulfate, 0.1 M sodium citrate, (v/v)dioxane, Нq 6.3. After 24 equilibration, crystal growth was induced by applying the seeding technique described in ref. 21, using crystals of 25 recombinant wild-type $Bet \ v \ 1$ as a source of seeds.

After about 4 months, crystals were harvested and analysed using X-rays generated from a Rigaku rotating anode as described in ref. 21 and the structure was solved using molecular replacement.

Structure of Bet v 1 (2628) mutant

The structural effect of the mutations was addressed by growing three-dimensional $Bet\ v\ 1\ (2628)$ protein crystals diffracting to 2.0 Å resolution when analysed by X-rays

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generated from a rotating anode. The substitutions Tyr5Val, Glu45Ser, Lys65Asn, Lys97Ser, Lys134Glu were verified by the Bet v 1 (2628) structure electron density map which also showed that the overall α -carbon backbone tertiary structure is preserved.

Structural analysis of Bet v 1 (2637) mutant

The structural integrity of the purified Bet v 1 (2637) analysed by circular dichroism mutant was spectroscopy. Figure 20 shows the CD spectra of recombinant Bet v 1.2801 (wildtype) and Bet v 1 (2637) mutant, recorded at close to equal concentrations. The overlap in peak amplitudes and positions in the CD spectra from the two recombinant proteins shows that the two preparations contain equal amounts of secondary structures strongly suggesting that the $\alpha\text{-carbon}$ backbone tertiary structure is not affected by the introduced amino acid substitutions.

IgE-binding properties of Bet v 1 (2628) and Bet v 1 (2637) mutants.

The IgE-binding properties of Bet v 1 (2628) and Bet v 1 (2637) as well as a 1:1 mix of Bet v 1 (2628) and Bet v 1 (2637) was compared with recombinant wild type Bet v 1.2801 in a fluid-phase IgE-inhibition assay using a pool of serum IgE derived from birch allergic patients.

As described in Example 1, recombinant Bet v 1.2801 was biotinylated at a molar ratio of 1:5 (Bet v 1 no. 2801:biotin). The inhibition assay was performed as follows: a serum sample (25 μ l) was incubated with solid phase anti IgE, washed, re-suspended and further incubated with a mixture of biotinylated Bet v 1.2801 and a given mutant or 1:1 mix of the two mutants. The amount

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of biotinylated Bet v 1.2801 bound to the solid phase was estimated from the measured RLU after incubation with acridinium ester labelled streptavidin. The degree of inhibition was calculated as the ratio between the RLU's obtained using buffer and mutant as inhibitor.

Figure 21 shows the inhibition of the binding of biotinylated recombinant Bet v 1.2801 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1.2801 and by Bet v 1 (2628), Bet v 1 (2637) and a 1:1 mix of Bet v 1 (2628) and Bet v 1 (2637).

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Bet v 1.2801 reaches 50% inhibition at about 5 ng whereas the corresponding concentration for Bet v 1 (2628) mutant is about 15-20 ng. This show that the point mutation introduced in the Bet v 1 (2628) mutant lowers the affinity for specific serum IgE by a factor of about 3-4.

The maximum level of inhibition reached by the Bet v 1 (2628) mutant protein is clearly lower compared to recombinant Bet v 1.2801. This may indicate that some of the specific IgE present in the serum pool are unable to recognise the Bet v 1 (2628) mutant protein due to the introduced point mutations.

30 Bet v 1 (2637) reaches 50% inhibition at about 400-500 ng showing that the point mutation introduced in the Bet v 1 (2637) mutant lowers the affinity for specific serum IgE by 80 to 100-fold compared to Bet v 1.2801. The large difference in IgE-binding is further supported by a clear difference in inclination of the inhibition curve obtained with Bet v 1(2637) mutant protein compared to

the inhibition curve for Bet v 1.2801. The different inclination provide evidence that the reduction in IgE-binding is due to a distinctly different epitope pattern of the mutant compared to Bet v 1.2801.

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In addition to the inhibition assays with single modified allergens a 1:1 mix of Bet 1 (2628) and Bet v 1 (2637) having same molar concentration of Bet v 1 as each of the samples with Bet 1 (2628) or Bet v 1 (2637), respectively was tested and showed full (100%) capacity to inhibit IgE-binding to rBet v 1.2801. The capacity to fully inhibit IgE-binding is a clear indication that all reactive epitopes present on Bet v 1.2801 were present in the 1:1 allergen mix. Further support comes from the comparable inclination of the two inhibition curves for Bet v 1.2801 and the allergen mix. Reduced IgE-reactivity of the mixed allergen sample is demonstrated by the need of a four-fold higher concentration of the allergen mix, when compared to Bet v 1.2801, for obtaining 50% inhibition of IgE-binding.

$\underline{\mathtt{T}}$ cell proliferation assay using mutated recombinant Bet v 1 allergens.

The analysis was carried out as described in ref. 15. Bet v 1 (2628) and Bet v 1(2637) mutant protein were both able to induce proliferation in T cell lines from birch pollen allergic patients with stimulation indices similar to recombinant and naturally occurring. This suggests that both of Bet v 1 (2628) and Bet v 1 (2637) mutant protein can each initiate the cellular immune response necessary for antibody production.

Histamine release assays with human basophil.

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Histamine release from basophil leucocytes was performed

as follows. Heparinized blood (20 ml) was drawn from each birch pollen patient, stored at room temperature, and used within 24 hours. Twenty-five microlitres heparinized whole blood was applied to glass fibre coated microtitre wells (Reference Laboratory, Denmark) and incubated with 25 microlitres of allergen or anti-IgE for 1 hour at 37°C. Thereafter the plates were rinsed and interfering substances were removed. Finally, histamine bound to the microfibres was spectrophotofluometrically.

Histamine release properties of Bet v 1 (2628) and Bet v 1 (2637) mutant protein.

Histamine release data is shown in Figure 22 and Figure 23. The potency of Bet v 1 (2628) and Bet v 1 (2637) mutant protein to induce histamine release in human basophil from two birch pollen allergic patients has been tested. In both cases the release curve of the mutated allergens to induce histamine release is clearly shifted to the right compared to the release curve of Bet v 1.2801. The shift indicate that the potency of Bet v 1 (2628) and Bet v 1 (2637) is reduced 3 to 10-fold.

Mutant Bet v 1 (2744) and mutant Bet v 1 (2753)

Bet v 1 (2744) and Bet v 1 (2753) was likewise constructed for use as a mixed allergen vaccine. In these mutated allergens point mutations were distributed in an all surface arranged fashion as shown in Figure 24 and Figure 25 and was again designed to affect different surface areas in the two molecules, respectively, as shown in Figure 26. However these modified allergens might individually be used as single allergen vaccines as well.

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Structural analysis of Bet v 1 (2744) mutant protein

The structural integrity of the purified Bet v 1 (2744) circular dichroism (CD) was analysed by mutant spectroscopy. the CD spectra Figure 27 shows recombinant Bet v 1.2801 (wildtype) and Bet v 1 (2744) mutant, recorded at close to equal concentrations. The overlap in peak amplitudes and positions in the CD spectra from the two recombinant proteins shows that the two preparations contain equal amounts of secondary structures strongly suggesting that the $\alpha\text{-carbon}$ backbone tertiary structure is not affected by the introduced amino acid substitutions.

15 Histamine release properties of Bet v 1 (2744)

Histamine release data from five experiments with basophil leucocytes from five different birch pollen allergic patients is shown in Figure 28 and Figure 29A-D. The potency of Bet v 1 (2744) mutant protein to induce histamine release in human basophil has been tested. The release curves of the mutated allergens are clearly shifted to the right compared to the release curve of Bet v 1.2801 indicating that the potency of Bet v 1 (2744) to release histamine is reduced 3 to 5-fold.

Mutant Bet v 1 (2733)

A Mutant Bet v 1 (2733) with nine primary mutations has expressed. recombinantly constructed and 30 been distribution of point mutations in Bet v 1 (2733) leave surface areas constituting >400Å² unaltered. several introduced point mutations Figure 30 show molecular surface of Bet v 1 (2733).

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This Example describes cloning of the gene encoding Der p 2 from *Dermatophagoides pteronyssinus* and construction of mutants with reduced IgE-binding affinity.

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PCR amplified products from first strand cDNA synthesis of Dermatophagoides pteronyssinus total RNA was obtained Wendy-Anne Smith and Dr. Wayne Thomas Telethon Institute for Child Health Research, 100 Roberts Rd, Subiaco, Western Australia 6008). During the amplification of the first strand cDNA library, Der p 2 had been selectively amplified using Der p 2 specific primers. PCR fragments were subsequently cloned into the England BioLabs). site of pUC19 (New sequencing of Der p 2 was performed using vector specific sense (5'-GGCGATTAAGTTGGGTAACGCCAGGG-3') and anti-sense (5'-GGAAACAGCTATGACCATGATTACGCC-3') primers.

A total of seven unique Der p 2 isoforms designated ALK-101, ALK-102, ALK-103, ALK-104, ALK-113, ALK-114, and ALK120 were identified. The clone entitled ALK-114 was chosen as starting point for generation of low-affinity IgE-mutants because of its high sequence identity with the Der p 2 NMR structure with the data base accession number 1A9V. Compared to ALK-114, the 6 other naturally occurring isoforms comprise the following substitutions:

ALK-101: M76V.

ALK-102: V40L, T47S.

ALK-103: M111L, D114N.

30 ALK-104: T47S, M111I, D114N.

ALK-113: T47S.

ALK-120: V40L, T47S, D114N.

Insertion of Der p 2 into $pGAPZ\alpha-A$

The gene encoding Der p 2 (ALK-114) was subsequently inserted into the pGAPZα-A vector (Invitrogen) secreted expression of Der p 2 in the yeast, Pichia pastoris. The gene was amplified using sense primer OB27 (5'-GGAATTCCTCGAGAAAAGAGATCAAGTCGATGTCAAAGATTGTGCC-3') anti-sense primer **OB28** (5'-CGTTCTAGACTATTAATCGCGGATTTTAGCATGAGTTGC-3') corresponding to the amino- and the carboxytermi of the Der p 2 polypeptide, respectively. The primers were extended in the 5'-end to accomodate the restriction sites Xho I and Xba I, respectively. The Xho I restriction site fuses the first codon of Der p 2 in frame with the nucleic acid sequence encoding the KEX2 cleavage site (LYS-ARG) $pGAPZ\alpha-A$. A single round of PCR amplification was performed in a 100 microliter (μ l) volume: template ALK-114 DNA, 1 X Expand polymerase buffer (available from Boehringer Mannheim), 0.2 millimolar (mM) each of the four dNTPs, 0.3 micromolar (μM) each of the sense and anti-sense primers and 2.5 Units of Expand polymerase (available from Boehringer Mannheim). The DNA amplified following 25 cycles of: 95°C seconds, 45°C for 30 seconds, 72°C for 1 minute, followed by 1 cycle of 72°C for 7 minutes. The resulting 475 base pair ALK-114 PCR fragment was purified using a QIAquick spin purification procedure (available from Qiagen). The purified DNA fragment was then digested with Xho I and Xba I, gel purified and ligated into similarly digested $pGAPZ\alpha-A$. The ligation reaction was trasformed E.coli strain DH5 α , resulting in plasmid, pCBo06.

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The nucleotide sequence of Der p 2 was confirmed by DNA sequencing before and after cloning and following in vitro mutagenesis (see below).

Der p 2 sequences

SEQ ID NO 1 corresponds to the nucleic acid sequence of Der p 2 (ALK-114):

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- 1 gatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaaagttttggtaccagga
- 51 tgccatggttcagaaccatgtatcattcatcgtggtaaaccattccaattggaagccgtt
- 121 ttcgaagccaaccaaaacacaaaaaccgctaaaattgaaatcaaagcctcaatcgatggt
- 181 ttagaagttgatgttcccggtatcgatccaaatgcatgccattacatgaaatgcccattg
- 10 241 gttaaaggacaacaatatgatattaaatatacatggaatgttccgaaaattgcaccaaaa
 - 301 tctgaaaatgttgtcgtcactgttaaagttatgggtgatgatggtgttttggcctgtgct
 - 361 attgctactcatgctaaaatccgcgattaa

SEQ ID NO 2 corresponds to the deduced amino acid sequence of Der p 2 (ALK-114):

- dqvdvkdcanheikkvlvpgchgsepciihrgkpfqleavfeangntktakieikasidq
- ${\tt 61} \qquad {\tt levdvpgidpnachymkcplvkgqqydikytwnvpkiapksenvvvtvkvmgddgvlaca}$
- 121 iathakird

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Insertion of pGAPZ α -A-Der p 2 into P. pastoris

The vector, pCBo06 was linearized using Avr II restriction enzyme and transformed into competent P. pastoris strain, X-33, as described in the Invitrogen manual. Recombinant cells resistant to 100 micrograms per milliliter (μ g/ml) of Zeocin were selected, and colony purifed on fresh YPD plates containing 100 μ g/ml Zeocin.

30 Expression and purification of recombinant Der p 2

A 250 ml of YPD medium (1% yeast extract, 2% peptone, 2% glucose) containing 100 μ g/ml Zeocin was inoculated with an overnight culture of recombinant yeast cells expressing Der p 2. The culture was grown at 30°C for 72 hours to obtain optimal Der p 2 expression. Cells were harvested by centrifugation and the resulting culture

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supernatant was saturated with 50% ammonium sulfate. Following centrifugation at 3000x g for 30 minutes, the supernatant was saturated with 80% ammonium sulfate. Following centrifugation, the pellet was resuspended in 150 millimolar (mM) NH₄HCO₃ and fractionated on a Superdex 75 gel filtration column, equilibrated with the same buffer. Der p 2 was eluted as a major peak corresponding to its expected molecular weight. The elution of Der p 2 was monitered both by SDS page electrophoresis, followed by silver staining and by immunoblot analysis using Der p 2 specific polyclonal antibodies.

Selection of amino acid residues for site-directed mutagenesis

Selection of amino acid residues for mutagenesis was based on identification of residues that are highly solvent exposed and highly conserved among allergens from House Dust Mites (Der p 2/f 2 and Eur m 2) and storage mites (Tyr p 2, Lep d 2, Gly d 2). Highly solvent exposed amino acid residues were identified visually by analysis of the molecular surface of the Der p 2 NMR structure (#1.9, 1A9V.pdb). Twelve amino acid residues were selected for mutagenesis: K6A, N10S, K15E, S24N, H30N, K48A, E62S, H74N, K77N, K82N, K100N and R128Q.

Site-directed mutagenesis

30 Construction of recombinant mutant allergens with single primary mutations and multiple combinations thereof, are described in the following.

Expression plasmids encoding Der p 2 mutants were produced using pCBo06 as DNA template. PCR reactions were

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sense and anti-sense primers performed using incorporating the specified mutations. Primer pairs used in the PCR reactions to generate the specified mutations are listed in Figure 31. The mutations are designated in bold and the restriction sites used in the subsequent cloning step are underlined in the figure. For the construction of mutants K6A, K15E, H30N, H74N and K82N, PCR reactions were performed essentially as described in the section "Cloning of Der p 2 into pGAPZ α -A". The purified PCR fragments were digested with the designated restriction enzyme sites (see Figure 31), gel purified, ligated into similarly digested pCBo06 and transformed into E.coli DH5 α .

The mutation E62S was generated using an alternative PCR 15 mutagenesis methodology described for the generation of Bet v 1 mutants in Example 1. Two mutation specific oligonucleotide primers were synthesized covering the specified mutations (OB47 and OB48, listed in Figure 31). additional primers used for the secondary 20 Two amplification step were OB27 and OB28 as described in the section: "Insertion of Der p 2 into pGAPZ α -A".

The mutant allergens produced are characterised using the same methods as described in example 4, e.g. circular dichroism (CD) spectroscopy, crystallisation, measurements of IgE binding properties, histamin-release, T-cell proliferation stimulation capacity, etc.

30 EXAMPLE 6

Mutated recombinant mite allergens (Der p 2) with improved safety for specific allergy vaccination

35 In this example the application of the concept of the

current invention on house dust mite allergens is exemplified by one allergen, Der p 2. Manipulation of other house dust mite allergens may be performed by equivalent procedures.

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Design of mutated recombinant Der p 2 molecules.

SEQ ID NO. 3 shows the nucleotide and deduced amino acid sequence of Der p 2-ALK-G clone, which is a wild type isoform.

SEQ ID NO. 3: Nucleotide and deduced amino acid sequence of Der p 2-ALK-G.

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	GAT D			AAA K	 _	 		 	 AAA K		45
20				TGC C			CCA P		CAT H	30	90
25									CAA Q	45	
	AAC N								GGT G	60	180
30									TAT Y	75	
									TAT Y	90	270
35									GTC V	105	
40		GTT V							GCT A	120	360
		 	 	AAA K	 					129	387

Fig. 32 shows a sequence alignment performed at the ExPaSy Molecular Biology Server (http://www.expasy.ch/) using the ClustalW algorithm on a BLAST search using the Der p 2-ALK-G amino acid sequence shown in SEQ ID NO. 3 as input sequence. The alignment includes sequences from house dust mite species, i.e. Der p 2, Der f 2 and Eur m 2. In Fig. 32 amino acid residues identical to amino

acids in the same position in the Der p 2-ALK-G protein sequence are highlighted using black letters on grey background. Non-identical amino acids are printed in black on a white background.

Surface structure images

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Amino acid sequences representing the house dust mite group 2 allergens have a similarity greater than 85 % and some of the molecular surface is conserved (grey-coloured zones), see Fig. 33.

Fig. 33 shows surface contours viewed from 4 different angles when superimposing the Der p 2-ALK-G protein sequence on to the published PDB:1A9V NMR structure, structure number 1 of 10 contained in the PDB file.

Conserved and highly solvent exposed amino acid spatially distributed over the entire surface within distances in the range of 25-30 Å are selected for mutation. In the sections below the following information is given: A list of amino acids considered to be appropriate for mutation (A), a list of the mutants designed (B) and the DNA sequences representing the mutants designed (C). Fig. 34 shows surface contours of mutant number 1 as an example. Grey colour indicates conserved amino acid residues. Black colour indicates amino acid residues selected for mutation.

30 A. List of amino acids selected for mutation

K15, S24, H30, R31, K48, E62, H74, K77, K82, K100, R128

B. List of mutants designed

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Mutant 1:

K15E, S24N, H30G, K48A, E62S, K77N, K82N, K100N

Mutant 2:

K15E, S24N, H30G, K48A, E62S, K77N, K82N, R128Q

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Mutant 3:

K15E, S24N, H30G, K48A, K77N, K82N, K100N, R128Q

Mutant 4:

10 K15E, S24N, H30G, E62S, K77N, K82N, K100N, R128Q

Mutant 5:

K15E, H30G, K48A, E62S, K77N, K82N, K100N, R128Q

15 Mutant 6:

S24N, H30G, K48A, E62S, K77N, K82N, K100N, R128Q

Mutant 7:

K15E, S24N, R31S, K48A, E62S, H74N, K82N, K100N

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Mutant 8:

K15E, S24N, R31S, K48A, E62S, H74N, K82N, R128Q

Mutant 9:

25 K15E, S24N, R31S, K48A, H74N, K82N, K100N, R128Q

Mutant 10:

K15E, S24N, R31S, E62S, H74N, K82N, K100N, R128Q

30 Mutant 11:

K15E, R31S, K48A, E62S, H74N, K82N, K100N, R128Q

Mutant 12:

S24N, R31S, K48A, E62S, H74N, K82N, K100N, R128Q

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C. Nucleotide sequence of mutants

Mutant 1:

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K15E, S24N, H30G, K48A, E62S, K77N, K82N, K100N

GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggtAACgaaccatgtatcattGGCcgtggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgccattatatg AACtgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaaatccgcgat

Mutant 2:

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K15E, S24N, H30G, K48A, E62S, K77N, K82N, R128Q

GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggtAACgaaccatgtatcattGGCcgtggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgccattatatg AACtgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaaaatctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGgat

Mutant 3:

K15E, S24N, H30G, K48A, K77N, K82N, K100N, R128Q

Mutant 4:

K15E, S24N, H30G, E62S, K77N, K82N, K100N, R128Q

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GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggtAACgaaccatgtatcattGGCcgtggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaaaaacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgccattatatg AACtgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGgat

Mutant 5:

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K15E, H30G, K48A, E62S, K77N, K82N, K100N, R128Q

GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggttcagaaccatgtatcattGGCcgtggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgccattatatg AACtgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGgat

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Mutant 6:

S24N, H30G, K48A, E62S, K77N, K82N, K100N, R128Q

Gatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaaagttttggtacca ggatgccatggtAACgaaccatgtatcattGGCcgtggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgccattatatg AACtgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGgat Mutant 7:

K15E, S24N, R31S, K48A, E62S, H74N, K82N, K100N

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GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggtAACgaaccatgtatcattcatAGCggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgcAACtatatg aaatgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatccgcgat

Mutant 8:

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K15E, S24N, R31S, K48A, E62S, H74N, K82N, R128Q

GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggtAACgaaccatgtatcattcatAGCggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgcAACtatatg aaatgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaaaatctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGgat

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Mutant 9:

K15E, S24N, R31S, K48A, H74N, K82N, K100N, R128Q

Mutant 10:

K15E, S24N, R31S, E62S, H74N, K82N, K100N, R128Q

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Mutant 11:

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K15E, R31S, K48A, E62S, H74N, K82N, K100N, R128Q

GatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaGAAgttttggtacca ggatgccatggttcagaaccatgtatcattcatAGCggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgcAACtatatg aaatgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGgat

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Mutant 12:

S24N, R31S, K48A, E62S, H74N, K82N, K100N, R128Q

Gatcaagtcgatgtcaaagattgtgccaatcatgaaatcaaaaagttttggtacca ggatgccatggtAACgaaccatgtatcattcatAGCggtaaaccattccaattggaa gctttattcgaagccaatcaaaactcaGCGacagctaaaattgaaatcaaagcttca atcgatggtttaAGCgttgatgttcccggtatcgatccaaatgcatgcAACtatatg aaatgtccattggttAACggacaacaatatgatattaaatatacatggaatgttcca 35 aaaattgcaccaAACtctgaaaatgttgtcgtcactgttaaagttttgggtgataat ggtgttttggcctgtgctattgctactcatgctaaaatcCAGqat

EXAMPLE 7

Mutated recombinant mite allergens (Der p 1) with improved safety for specific allergy vaccination

In this example the application of the concept of the current invention on house dust mite allergens is exemplified by one allergen, Der p 1. Manipulation of other house dust mite allergens may be performed by equivalent procedures.

Design of mutated recombinant Der p 1 molecules.

SEQ ID NO. 4 shows the nucleotide and deduced amino acid sequence of Der p 1-ALK clone, which is a wild-type isoform.

SEQ ID NO. 4: Nucleotide and deduced amino acid sequence 20 of Der p 1-ALK

25	ACT T	AAC N	GCC A	TGC C	AGT S	ATC I	AAT N	GGA G	AAT N	GCT A	CCA P	GCT A	GAA E	ATC I	GAT D		45
	TTG L	CGA R	CAA Q	ATG M	CGA R	ACT T	GTC V	ACT T	CCC P	ATT I	CGT R	ATG M	CAA Q	GGA G	GGC G		90
30	TGT C	GGT G	TCA S	TGT C	TGG W	GCT A	TTC F	TCT S	GGT G	GTT V	GCC A	GCA A	ACT T	GAA E	TCA S		135
35	GCT A	TAT Y	TTG L	GCT A	TAC Y	CGT R	AAT N	CAA Q	TCA S	TTG L	GAT D	CTT L	GCT A	GAA E	CAA Q		180
	GAA E	TTA L	GTC V	GAT D	TGT C	GCT A	TCC S	CAA Q	CAC H	GGT G	TGT C	CAT H	GGT G	GAT D	ACC T	75	225
40	ATT I	CCA P	CGT R	GGT G	ATT I	GAA E	TAC Y	ATC I	CAA Q	CAT H	AAT N	GGT G	GTC V	GTC V	CAA Q	90	270
	GAA E	AGC S	TAC Y	TAT Y	CGA R	TAC Y	GTT V	GCA A	CGA R	GAA E	CAA Q	TCA S	TGC C	CGA R	CGA R	105	
45	CCA P	AAT N	GCA A	CAA Q	CGT R	TTC F	GGT G	ATC I	TCA S	AAC N	TAT Y	TGC C	CAA Q	ATT I	TAC Y	120	
50	CCA P	CCA P	AAT N	GTA V	AAC N	AAA K	ATT I	CGT R	GAA E	GCT A	TTG L	GCT A	CAA Q	ACC T	CAC H	135	
	AGC S	GCT A	ATT I	GCC A	GTC V	ATT I	ATT I	GGC G	ATC I	AAA K	GAT D	TTA L	GAC D	GCA A	TTC F	150	
55	CGT R	CAT H	TAT Y	GAT D	GGC G	CGA R	ACA T	ATC I	ATT I	CAA Q	CGC R	GAT D	AAT N	GGT G	TAC Y	165	

CAA CCA AAC TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GCA 540 Q P N Y H A V N I V G Y S N A CAA GGT GTC GAT TAT TGG ATC GTA CGA AAC AGT TGG GAT ACC AAT 585 5 YWIVRNS D TGG GGT GAT AAT GGT TAC GGT TAT TTT GCT GCC AAC ATC GAT TTG 630 G D N G Y G Y F A A N I D L 210 10 ATG ATG ATT GAA GAA TAT CCA TAT GTT GTC ATT CTC 666 MMIEEYPYV 222

Fig. 35 shows a sequence alignment performed at the ExPaSy Molecular Biology Server (http://www.expasy.ch/) using the ClustalW algorithm on a BLAST search using the Der p 1-ALK amino acid sequence shown in SEQ ID NO. 4 as input sequence. The alignment includes sequences from house dust mite species, i.e. Der p 1, Der f 1 and Eur m 1. In Fig. 35 amino acid residues identical to amino acids in the same position in the Der p 1-ALK protein sequence are highlighted using black letters on grey background. Non-identical amino acids are printed in black on a white background.

Surface structure images

Amino acid sequences representing the house dust mite group 1 allergens are similar to a certain degree and some of the molecular surface is conserved (grey-coloured zones), see Fig. 36. Fig. 36 shows surface contours viewed from 4 different angles when superimposing the Der p 1-ALK protein sequence on to a Der p 1 molecular structure model.

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Conserved and highly solvent exposed amino acid spatially distributed over the entire surface within distances in the range of 25-30 Å are selected for mutation. In the sections below the following information is given: A list of amino acids considered to be appropriate for mutation (A), a list of the mutants designed (B) and the DNA sequences representing the mutants designed (C). Fig. 37

shows surface contours of mutant number 11 as an example. Grey colour indicates conserved amino acid residues. Black colour indicates amino acid residues selected for mutation.

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A. List of amino acids selected for mutation

E13, P24, R20, Y50, S67, R78, R99, Q109, R128, R156, R161, P167, W192

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B. List of mutants designed

Mutant 1:

P24T, Y50V, R78E, R99Q, R156Q, R161E, P167T

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Mutant 2:

P24T, Y50V, R78Q, R99E, R156E, R161Q, P167T

Mutant 3:

20 R20E, Y50V, R78Q, R99Q, R156E, R161E, P167T

Mutant 4:

R20Q, Y50V, R78E, R99E, R156Q, R161Q, P167T

25 Mutant 5:

P24T, Y50V, S67N, R99E, R156Q, R161Q, P167T

Mutant 6:

R20E, Y50V, S67N, R99E, R156Q, R161E, P167T

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Mutant 7:

R20Q, Y50V, S67N, R99Q, R156E, R161E, P167T

Mutant 8:

35 E13S, P24T, Y50V, R78E, R99Q, Q109D, R128E, R156Q, R161E, P167T

Mutant 9:

E13S, P24T, Y50V, R78Q, R99E, Q109D, R128Q, R156E, R161Q, P167T

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Mutant 10:

E13S, P24T, Y50V, R78E, R99Q, Q109D, R128E, R156Q, R161E, P167T, W192F

10 Mutant 11:

E13S, P24T, Y50V, R78Q, R99E, Q109D, R128Q, R156E, R161Q, P167T, W192F

C. Nucleotide sequences of mutants

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Mutant 1:

P24T, Y50V, R78E, R99Q, R156Q, R161E, P167T

20 ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT GAA ATC GAT TTG CGA CAA ATG CGA 60 ACT GTC ACT ACC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120 GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180 GAA TTA GTC GAT TGT GCT TCC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA GAA GGT ATT 240 GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA CAG GAA 300 25 CAA TCA TGC CGA CGA AAT GCA CAA CGT TTC GGT ATC TCA AAC TAT TGC CAA ATT TAC 360 CCA CCA AAT GTA AAC AAA ATT CGT GAA GCT TTG GCT CAA ACC CAC AGC GCT ATT GCC GTC 420 ATT ATT GGC ATC AAA GAT TTA GAC GCA TTC CGT CAT TAT GAT GGC CAG ACA ATC ATT CAA 480 GAA GAT AAT GGT TAC CAA ACC AAC TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GCA 540 CAA GGT GTC GAT TAT TGG ATC GTA CGA AAC AGT TGG GAT ACC AAT TGG GGT GAT AAT GGT 600 30 TAC GGT TAT TTT GCT GCC AAC ATC GAT TTG ATG ATG ATT GAA GAA TAT CCA TAT GTT GTC 660 ATT CTC 666

Mutant 2:

35 P24T, Y50V, R78Q, R99E, R156E, R161Q, P167T

Mutant 3:

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R20E, Y50V, R78Q, R99Q, R156E, R161E, P167T

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Mutant 4:

R20Q, Y50V, R78E, R99E, R156Q, R161Q, P167T

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT GAA ATC GAT TTG CGA CAA ATG CAG 60
ACT GTC ACT CCC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120

Mutant 5:

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P24T, Y50V, S67N, R99E, R156Q, R161Q, P167T

Mutant 6:

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R20E, Y50V, S67N, R99E, R156Q, R161E, P167T

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT GAA ATC GAT TTG CGA CAA ATG GAA 60

ACT GTC ACT CCC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120

GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180

GAA TTA GTC GAT TGT GCT AAC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA CGT GGT ATT 240

GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA GAA GAA 300
CAA TCA TGC CGA CGA CCA AAT GCA CAA CGT TTC GGT ATC TCA AAC TAT TGC CAA ATT TAC 360
CCA CCA AAT GTA AAC AAA ATT CGT GAA GCT TTG GCT CAA ACC CAC AGC GCT ATT GCC GTC 420
ATT ATT GGC ATC AAA GAT TTA GAC GCA TTC CGT CAT TAT GAT GGC CAG ACA ATC ATT CAA 480
GAA GAT AAT GGT TAC CAA ACC AAC TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GGT 600
TAC GGT TAT TTT GCT GCC AAC ATC GAT TTG ATG ATG ATT GAA GAA TAT CCA TAT GTT GTC 660
ATT CTC

10 Mutant 7:

R20Q, Y50V, S67N, R99Q, R156E, R161E, P167T

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT GAA ATC GAT TTG CGA CAA ATG CAG 60 15 ACT GTC ACT CCC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120 GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180 GAA TTA GTC GAT TGT GCT AAC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA CGT GGT ATT 240 GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA CAG GAA 300 CAA TCA TGC CGA CGA CCA AAT GCA CAA CGT TTC GGT ATC TCA AAC TAT TGC CAA ATT TAC 360 20 CCA CCA AAT GTA AAC AAA ATT CGT GAA GCT TTG GCT CAA ACC CAC AGC GCT ATT GCC GTC 420 ATT ATT GGC ATC AAA GAT TTA GAC GCA TTC CGT CAT TAT GAT GGC GAA ACA ATC ATT CAA 480 GAA GAT AAT GGT TAC CAA ACC AAC TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GCA 540 CAA GGT GTC GAT TAT TGG ATC GTA CGA AAC AGT TGG GAT ACC AAT TGG GGT GAT AAT GGT 600 TAC GGT TAT TTT GCT GCC AAC ATC GAT TTG ATG ATG ATT GAA GAA TAT CCA TAT GTT GTC 660 25 ATT CTC 666

Mutant 8:

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E13S, P24T, Y50V, R78E, R99Q, Q109D, R128E, R156Q, R161E, 30 P167T

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT AGC ATC GAT TTG CGA CAA ATG CGA 60

ACT GTC ACT ACC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120

GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180

GAA TTA GTC GAT TGT GCT TCC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA GAA GGT ATT 240

GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA CAG GAA 300

CAA TCA TGC CGA CGA CCA AAT GCA GAT CGT TTC GGT ATC TCA AAC TAT TGC CAA ATT TAC 360

CCA CCA AAT GTA AAC AAA ATT GAA GAA GCT TTG GCT CAA ACC CAC AGC GCT ATT GCC GTC 420

ATT ATT GGC ATC AAA GAT TTA GAC GCA TTC CGT CAT TAT GAT GGC CAG ACA ATC ATT CAA 480

GAA GAT AAT GGT TAC CAA ACC AAC TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GCA 540

CAA GGT GTC GAT TAT TGG ATC GTA CGA AAC AGT TTG ATG ATG ATT GAA GAA TAT CCA TAT GTT GTC 660

ATT CTC

Mutant 9:

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E13S, P24T, Y50V, R78Q, R99E, Q109D, R128Q, R156E, R161Q, P167T

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT AGC ATC GAT TTG CGA CAA ATG CGA 60 15 ACT GTC ACT ACC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120 GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180 GAA TTA GTC GAT TGT GCT TCC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA CAG GGT ATT 240 GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA GAA GAA 300 CAA TCA TGC CGA CGA CCA AAT GCA GAT CGT TTC GGT ATC TCA AAC TAT TGC CAA ATT TAC 360 20 CCA CCA AAT GTA AAC AAA ATT CAG GAA GCT TTG GCT CAA ACC CAC AGC GCT ATT GCC GTC 420 ATT ATT GGC ATC AAA GAT TTA GAC GCA TTC CGT CAT TAT GAT GGC GAA ACA ATC ATT CAA 480 CAG GAT AAT GGT TAC CAA ACC AAC TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GCA 540 CAA GGT GTC GAT TAT TGG ATC GTA CGA AAC AGT TGG GAT ACC AAT TGG GGT GAT AAT GGT 600 TAC GGT TAT TTT GCT GCC AAC ATC GAT TTG ATG ATG ATT GAA GAA TAT CCA TAT GTT GTC 660 25 ATT CTC 666

Mutant 10:

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E13S, P24T, Y50V, R78E, R99Q, Q109D, R128E, R156Q, R161E, 30 P167T, W192F

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT AGC ATC GAT TTG CGA CAA ATG CGA 60

ACT GTC ACT ACC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120

GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180

GAA TTA GTC GAT TGT GCT TCC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA GAA GGT ATT 240

GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA CAG GAA 300

Mutant 11:

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E13S, P24T, Y50V, R78Q, R99E, Q109D, R128Q, R156E, R161Q, P167T, W192F

ACT AAC GCC TGC AGT ATC AAT GGA AAT GCT CCA GCT AGC ATC GAT TTG CGA CAA ATG CGA 60

ACT GTC ACT ACC ATT CGT ATG CAA GGA GGC TGT GGT TCA TGT TGG GCT TTC TCT GGT GTT 120

GCC GCA ACT GAA TCA GCT TAT TTG GCT GTG CGT AAT CAA TCA TTG GAT CTT GCT GAA CAA 180

GAA TTA GTC GAT TGT GCT TCC CAA CAC GGT TGT CAT GGT GAT ACC ATT CCA CAG GGT ATT 240

GAA TAC ATC CAA CAT AAT GGT GTC GTC CAA GAA AGC TAC TAT CGA TAC GTT GCA GAA GAA 300

CAA TCA TGC CGA CGA CCA AAT GCA GAT CGT TTC GGT ATC TCA AAC TAT TGC CAA ATT TAC 360

CCA CCA AAT GTA AAC AAA ATT CAG GAA GCT TTG GCT CAA ACC CAC AGC GCT ATT GCC GTC 420

ATT ATT GGC ATC AAA GAT TTA GAC GCA TAT CAC GCT GTC AAC ATT GTT GGT TAC AGT AAC GCA 540

CAA GGT GTC GAT TAT TGG ATC GTA CGA AAC AGT TTT GAT ACC AAT TGG GGT GAT AAT GGT 600

TAC GGT TAT TTT GCT GCC AAC ATC GAT TTG ATG ATG ATT GAA GAA TAT CCA TAT GTC GTC 660

ATT CTC

EXAMPLE 8

Mutated recombinant grass allergens (Phl p 5) with improved safety for specific allergy vaccination

In this example the application of the concept of the current invention on grass pollen allergens is exemplified by one allergen, Phl p 5. Manipulation of other grass pollen allergens may be performed by equivalent procedures.

Design of mutated recombinant Phl p 5 molecules

SEQ ID NO. 5 shows the nucleotide and deduced amino acid sequence of the Phl p 5.0103 clone, which is a wild-type isoform.

SEQ ID NO. 5: Nucleotide and deduced amino acid sequence of Phl p $5.0103\,.$

gcc	gat	ctc	ggt	tac	ggc	ccc	gcc	acc	:cca	gct	.gcc	ccg	gcc	gcc	ggc	tac	acc	ccc	gcc	60
Α	D	L	G	Y	G	P	A	Т	P	A	A	P	A	A	G	Y	т	P	A	20
acc	ccc	gcc	gcc	ccg	gcc	gga	gcg	gag	rcca	gca	ggt	aag	ıgcg	acg	acc	gag	gag	cag	aag	120
т	P	A	Α	P	Α	G	A	E	P	Α	G	K	A	Т	т	E	E	Q	K	40
ctg	atc	gag	aag	atc	aac	gcc	ggc	ttc	aag	gcg	gcc	ttg	gcc	gct	gcc	gcc	ggc	gtc	ccg	180
L	I	E	K	I	N	A	G	F	K	Α	A	L	Α	Α	A	A	G	v	P	60
cca	gcg	gac	aag	tac	agg	acg	ttc	gto	gca	acc	ttc	ggc	gcg	gcc	tcc	aac	aag	gcc	ttc	240
P	A	D	K	Y	R	T	F	V	Α	Т	F	G	Α	Α	s	N	K	Α	F	80
gcg	gag	ggc	ctc	tcg	ggc	gag	ccc	aag	ggc	gcc	gcc	gaa	tcc	ago	tcc	aag	gcc	gcg	ctc	300
A	E	G	L	s	G	E	P	K	G	Α	Α	E	s	s	S	K	Α	Α	L	100
acc	tcc	aag	ctc	gac	gcc	gcc	tac	aag	ctc	gcc	tac	aag	aca	gcc	gag	ggc	gcg	acg	cct	360
T	s	K	L	D	Α	Α	Y	K	L	A	Y	K	T	A	E	G	Α	Т	P	120
gag	gcc	aag	tac	gac	gcc	tac	gtc	gcc	acc	gta	ago	gag	gcg	ctc	cgc	atc	atc	gcc	ggc	420
E	A	K	Y	D	A	Y	V	A	Т	V	s	E	Α	L	R	I	I	A	G	140
acc	ctc	gag	gtc	cac	gcc	gtc	aag	ccc	gcg	gcc	gag	gag	gto	aag	gtc	atc	ccc	gcc	ggc	480
Т	L	E	V	Н	Α	V	K	P	A	A	E	E	V	K	V	Ţ	P	Α	G	160
jag	ctg	cag	gtc	atc	gag	aag	gtc	gac	gcc	gcc	ttc	aag	gtc	gct	gcc	acc	gcc	gcc	aac	540
Ε	L	Q	V	I	E	K	V	D	A	A	F	K	V	A	A	Т	A	A	N	180
jcc	gcc	CCC	gcc	aac	gac	aag	ttc	acc	gtc	ttc	gag	gcc	gcc	ttc	aac	gac	gcc	atc	aag	600
A	Α	P	A	N	D	K	F	Т	V	F	Ε	A	A	F	N	D	Α	I	K	200
gcg	agc	acg	ggc	ggc	gcc	tac	gag	agc	tac	aag	ttc	atc	ccc	gcc	ctg	gag	gcc	gcc	gtc	660
A	S	Т	G	G	Α	Y	E	S	Y	K	F	I	P	A	L	E	A	A	V	220
aag	cag	gcc	tac	gcc	gcc	acc	gtc	gcc	acc	gcg	ccg	gag	gtc	aag	tac	act	gtc	ttt	gag	720
-	Q		_				V		-	A	_	E	-	K		_	٠	F	Ε	240
acc	gca	ctg	aaa															CCC	gcc	780
Т	A	L	K	K	Α	Ι	Т	A	M	S	E	Α	Q	K	А	A	K	P	Α	260
gccgctgccaccgccaccgccaccgccgttggcgcggccaccggcgccaccgcc								gcc	840											
	A		_		Т		Т	A	Α	V	G	A	Α	T	G	Α	Α	Т	Α	280
зсt	act	ggt	ggc	tac	aaa	gtc														861

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Fig. 38 shows a sequence alignment performed at the ExPaSy Molecular Biology Server (http://www.expasy.ch/) using the ClustalW algorithm on a BLAST search using the Phl p 5.0103 amino acid sequence shown in SEQ ID NO. 5 as input sequence. The alignment includes group 5 allergen sequences from grass species, i.e. Phl p 5, Poa p 5, Lol p 5, Hol l 5, Pha a 5, Hor v 9 and Hor v 5. In Fig. 38 amino acid residues identical to amino acids in the same position in the Phl p 5.0103 protein sequence are highlighted using black letters on grey background. Non-identical amino acids are printed in black on a white background.

15 Surface structure images

Amino acid sequences representing the grass pollen group 5 allergens are similar to a certain degree and some of the molecular surface is conserved (grey-coloured zones), see Fig. 39. Fig. 39 shows surface contours viewed from 4 different angles when superimposing the Phl p 5.0103 protein sequence on to a Phl p 5 molecular structure model. The structure model encompass the molecule in two halves, Model A (amino acid 34-142) shown in Fig. 39A, and Model B (amino acid 149-259) shown in Fig. 39B.

Highly solvent exposed amino acid spatially distributed over the entire surface within distances in the range of 25-30 Å are selected for mutation. In the sections below, the following information is given: A list of amino acids considered to be appropriate for mutation (A), a list of designed (B) the mutants and DNA representing the mutants designed (C). Fig. 40 A and B shows surface contours of mutant number 1 Model A and Model В, respectively, as an example. Grey colour indicates conserved amino acid residues. Black colour

indicates amino acid residues selected for mutation.

A: List of amino acids selected for mutation

5 I45, R66, E133, R136, I137, D186, F188, K211, P214, Q222, P232, L243, Q254

B. List of mutants designed

10 Mutant 1:

I45K, E133S, F188I, Q222K, L243E, Q254K

Mutant 2:

R66N, E133S, F188I, Q222K, L243E, Q254K

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Mutant 3:

I45K, R136S, F188I, Q222K, L243E, Q254K

Mutant 4:

20 I45K, I137K, F188I, Q222K, L243E, Q254K

Mutant 5:

I45K, E133S, D186H, Q222K, L243E, Q254K

25 Mutant 6:

I45K, E133S, Q222K, P232G, L243E, Q254K

Mutant 7:

I45K, E133S, F188I, P214G, L243E, Q254K

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Mutant 8:

I45K, E133S, F188I, K211N, L243E, Q254K

Mutant 9:

35 R66N, R136S, F188I, Q222K, L243E, Q254K

Mutant 10: R66N, I137K, F188I, Q222K, L243E, Q254K Mutant 11: 5 I45K, E133S, D186H, P214G, L243E, Q254K Mutant 12: I45K, E133S, D186H, K211N, L243E, Q254K Mutant 13: 10 I45K, E133S, P214G, P232G, L243E, Q254K Mutant 14: I45K, E133S, K211N, P232G, L243E, Q254K 15 C. Nucleotide sequence of mutants Mutant 1: 20 I45K, E133S, F188I, Q222K, L243E, Q254K: geogateteggttacggccccgccaccccaqctqcccggccgccqqctacacccccqcc 60 acccccgccgcccggccggagcgagccagcaggtaaggcgacgaccgaggagcagaag 120 180 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccgcggcgtcccg 25 $\verb|ccagcggacaagtacaggacgttcgtcgcaaccttcggcgggcctccaacaaggccttc|$ 240 geggagggetetegggegageeeaagggegeegegaateeageteeaaggeegegete 300 360 acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggcgcgacgcct 420 gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc accetegaggtecacgecgteaagcecgeggecgaggaggteaaggteatccecgeegge 480 30 gagctgcaggtcatcgagaaggtcgacgccgccttcaaggtcgctgccaccgccgccaac 540 gccgccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag 600 660 gcgagcacgggcgccctacgagagctacaagttcatccccgccctggaggccgccgtc 720 aagAAAgcctacgccgccaccgtcgccaccgcgccggaggtcaagtacactgtctttgag accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc 780 35 geegetgecacegecacegeacegeegttggegegecaceggegecacegee 840 gctactggtggctacaaagtc 861 Mutant 2: 40 R66N, E133S, F188I, Q222K, L243E, Q254K: 60 gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc acccccgccgcccggccggagcggagccagcaggtaaggcgacgacgacgaggagcagaag 120

ctgatcgagaagatcaacgccggcttcaaggcggccttggccgctgccggcggccccg

 $\verb|ccagcggacaagtacAACacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc|\\$

180

geggagggetetegggegageceaagggegeegeegaateeageteeaaggeegegete 300 360 gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc 420 ${\tt accctcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc}$ 480 gagctgcaggtcatcgagaaggtcgacgccgccttcaaggtcgctgccaccgccgccaac 540 $\verb|gccgccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag|$ 600 gcgagcacgggcggcgcctacgagagctacaagttcatccccgccctggaggccgccgtc 660 aagAAAgcctacgccgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag 720 $\verb"accgcaGAA" a \verb"aaaa" aggccatcaccgccatgtccgaagcaAAA aaggctgccaagcccgcc$ 780 10 $\tt gccgctgccaccgccaccgccaccgccgcttggcgcgccaccggcgccaccgcc$ 840 gctactggtggctacaaagtc 861

Mutant 3:

15 I45K, R136S, F188I, Q222K, L243E, Q254K:

	gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc	60
	acccccgccgcccggccggagcgagccagcaggtaaggcgacgaccgaggagcagaag	120
20	ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccgccggcgtcccg	180
	ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc	240
	gcggagggcctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc	300
	acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggcgcgacgcct	360
	gaggccaagtacgacgcctacgtcgccaccgtaagcgaggcgctcAGCatcatcgccggc	420
	accetcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc	480
25	gagctgcaggtcatcgagaaggtcgacgccgccttcaaggtcgctgccaccgccgccaac	540
	gccgccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag	600
	gcgagcacgggcggcctacgagagctacaagttcatccccgccctggaggccgccgtc	660
	aagAAAgcctacgccgccaccgtcgccaccgcgccggaggtcaagtacactgtctttgag	720
	accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc	780
30	gccgctgccaccgccaccgcaaccgccgccgttggcgcggccaccggcgccaccgcc	840
	gctactggtggctacaaagtc	861

Mutant 4:

35 I45K, I137K, F188I, Q222K, L243E, Q254K:

40	gccgatctcggttacggcccgccaccccagctgccccggccgcggctacacccccgcc accccgccgcccggccggagcggag	60 120 180 240 300
	gcggagggcctctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc acctccaagctcgacgcctacaagctcgcctacaagacagccgagggcgcgacgcct gaggccaagtacgacgcctacgtcgccaccgtaagcgaggcgctccgcAAAatcgccggc	360 420
45	accttcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc gagctgcaggtcatcgagaaggtcgacgccgcttcaaggtcgctgccaccgccgccaac gccgcccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag gcgagcacggggggcgcctacgagagctacaagttcatccccgccctggaggccgccgtc	480 540 600 660
50	aagAAAgcctacgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagccgcc gccgctgccaccgccaccgcaaccgccgttggcgcggccaccggcgccaccgcc gctactggtggctacaaagtc	720 780 840 861

Mutant 5:

55 I45K, E133S, D186H, Q222K, L243E, Q254K:

gccgatctcggttacggccccgccaccccagctgccccggccgccgccgctacacccccgcc 60 acccccgccgcccggccggagccgagccagcaggtaaggcgacgaccgaggagcagaag 120 180 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccgccggcgtcccg $\verb|ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc|$ 240 geggagggetetegggegageceaagggegeegeegaateeageteeaaggeegegete 300 360 $\verb|acctccaagctcgacgcctacaagctcgcctacaagacagccgagggcgccacgcct|$ $\tt gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc$ 420 480 accetegaggtecacgecgteaagceegeggeegaggaggteaaggteateceegeegge gagetgcaggtcategagaaggtegaegccgcettcaaggtegetgccacegccgaca 540 10 600 gccgccccgccaacCATaagttcaccgtcttcgaggccgccttcaacgacgccatcaag $\tt gcgagcacggcgcctacgagagctacaagttcatccccgccctggaggccgccgtc$ 660 aagAAAgcctacgccgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag 720 780 accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc $\tt gccgctgccaccgccaccgccaccgccgccgttggcgcggccaccggcgccaccgcc$ 840 15 gctactggtggctacaaagtc 861

Mutant 6:

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I45K, E133S, Q222K, P232G, L243E, Q254K:

gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc 60 acccccgccgcccggagcggagccagcaggtaaggcgacgaccgaggagcagaag 120 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccggcggtcccg 180 240 ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc 300 gcggagggcctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc acctccaagetcgacgcctacaagetcgcctacaagacagccgagggcgcgacgcct 360 420 gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc accctcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc 480 gagetgeaggteategagaaggtegaegeetteaaggtegetgeeaeegeegeeaae 540 $\verb"gccgccccgccaacgacaagttcaccgtcttcgaggccgccttcaacgacgccatcaag"$ 600 gcgagcacgggcggcctacgagagctacaagttcatccccgccctggaggccgccgtc 660 aagAAAgcctacgccgccaccgtcgccaccgcgGGCgaggtcaagtacactgtctttgag 720 accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc 780 840 gccgctgccaccgccaccgccgccgttggcgcgccaccggcgccaccgcc gctactggtggctacaaagtc 861

Mutant 7:

I45K, E133S, F188I, P214G, L243E, Q254K:

40 60 gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc acccccgccgcccggccggagccggccagcaggtaaggcgacgaccgaggagcagaag 120 180 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccggcgtcccg 240 ccagcggacaagtacaggacgttcgtcgcaaccttcggcgggcctccaacaaggccttc 45 gcggagggcctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc 300 acctccaagetcgacgcctacaagetcgcctacaagacagecgagggcgcgacgcct 360 420 gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc accctcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc 480 gagctgcaggtcatcgagaaggtcgacgccgccttcaaggtcgctgccaccgccgaccaac 540 50 600 gccgccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag $\tt gcgagcacggcgcctacgagagctacaagttcatcGGCgccctggaggccgccgtc$ 660 720 aagcaggcctacgccgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag 780 accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc 840 gccgctgccaccgccaccgcaaccgccgttggcgcggccaccggcgccaccgcc 55 gctactggtggctacaaagtc 861

Mutant 8:

I45K, E133S, F188I, K211N, L243E, Q254K:

	gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc	60
5	acccccgccgcccggccggagcgagccagcaggtaaggcgacgaccgaggagcagaag	120
	ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccgccggcgtcccg	180
	ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc	240
	gcggagggcctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc	300
	acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggcgcgacgcct	360
10	gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc	420
	accetegaggtecacgecgteaagcecgeggeegaggaggteaaggteateceegeegge	480
	gagetgeaggteategagaaggtegacgeegtteaaggtegetgeeacegeegeeaac	540
	gccgccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag	600
	gcgagcacgggcgcctacgagagctacAACttcatccccgccctggaggccgccgtc	660
15	aagcaggcctacgccgccaccgtcgccaccgcgccggaggtcaagtacactgtctttgag	720
	accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc	780
	gccgctgccaccgccaccgccaccgccgttggcgcggccaccggcgccaccgcc	840
	gctactggtggctacaaagtc	861

20 Mutant 9:

R66N, R136S, F188I, Q222K, L243E, Q254K:

	gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc	60
25	acccccgccgcccggccggagcgagccagcaggtaaggcgacgaccgaggagcagaag	120
	ctgatcgagaagatcaacgccggcttcaaggcggccttggccgctgccggcggccccg	180
	ccagcggacaagtacAACacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc	240
	gcggagggcctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc	300
	acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggggggcgccct	360
30	gaggccaagtacgacgcctacgtcgccaccgtaagcgaggcgctcAGCatcatcgccggc	420
	accetegaggtecacgeegteaageeegeggeegaggaggteaaggteateeeegeegge	480
	gagetgeaggteategagaaggtegaegeegteteaaggtegetgeeacegeegeeaac	540
	gccgcccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag	600
	gcgagcacgggcggcgcctacgagagctacaagttcatccccgccctggaggccgccgtc	660
35	aagAAAgcctacgccgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag	720
	accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc	780
	gccgctgccaccgccaccgccaccgccgttggcgcggccaccggcgccaccgcc	840
	gctactggtggctacaaagtc	861

40 Mutant 10:

R66N, I137K, F188I, Q222K, L243E, Q254K:

	gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc	60
45	acccccgccgcccggccggagcggagccagcaggtaaggcgacgaccgaggagcagaag	120
	ctgatcgagaagatcaacgccggcttcaaggcggccttggccgctgccgccggcgtcccg	180
	ccagcggacaagtacAACacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc	240
	geggagggetetegggegageceaagggegeegeaateeageteeaaggeegete	300
	acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggcgcgacgcct	360
50	gaggccaagtacgacgcctacgtcgccaccgtaagcgaggcgctccgcAAAatcgccggc	420
	acctcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc	480
	gagctgcaggtcatcgagaaggtcgacgccgccttcaaggtcgctgccaccgccgccaac	540
	gccgccccgccaacgacaagATTaccgtcttcgaggccgccttcaacgacgccatcaag	600
	gcgagcacgggcgcctacgagagctacaagttcatccccgccctggaggccgccgtc	660
55	aagAAAgcctacgccgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag	720
	accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc	780
	gcgctgccaccgcaccgcaccgccgttggcggcaccggcgcaccgc	840

861 gctactggtggctacaaagtc Mutant 11: 5 I45K, E133S, D186H, P214G, L243E, Q254K: 60 gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc acccccgccgcccggaccggagccagcaggtaaggcgacgaccgaggagcagaag 120 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccgccggcgtcccg 180 10 $\verb|ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc|$ 240 300 gcggaggqcctctcggqcqaqcccaagqqcqccqccqaatccaqctccaaqqccqcqctc acctccaagctcgacgcctacaagctcgcctacaagacagccgagggcgcgacgcct 360 420 gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc accetegaggtecacgccgtcaagcccgcgggccgaggaggtcaaggtcatccccgccggc 480 15 gagetgcaggtcatcgagaaggtcgacgccttcaaggtcgctgccaccgccgaca 540 gccgccccgccaacCATaagttcaccgtcttcgaggccgccttcaacgacgccatcaag 600 $\tt gcgagcacggcgccctacgagagctacaagttcatcGGCgccctggaggccgccgtc$ 660 ${\tt aagcaggcctacgccgccaccgtcgccaccgcgccggaggtcaagtacactgtctttgag}$ 720 accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc 780 20 gccgctgccaccgccaccgcaccgccgttggcgcggccaccggcgccaccgcc 840 gctactggtggctacaaagtc 861 Mutant 12: 25 I45K, E133S, D186H, K211N, L243E, Q254K: $\tt gccgatctcggttacggccccgccaccccagctgccccggccgccgccacccccgcc$ 60 acccccgccgcccggaccggagccagcaggtaaggcgacgaccgaggagcagaag 120 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccgccggcgtcccg 180 ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc 240 300 geggagggeetetegggegageeeaagggegeegegaateeageteeaaggeegegete ${\tt acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggcgcgacgcct}$ 360 gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc 420 accctcgaggtccacgccgtcaagcccgcggccgaggaggtcaaggtcatccccgccggc 480 35 gagetgeaggteategagaaggtegaegeetteaaggtegetgeeaeegeegeeaae 540 gccgccccgccaacCATaagttcaccgtcttcgaggccgccttcaacgacgccatcaag 600 gcgagcacgggcgcctacgagagctacAACttcatccccgccctggaggccgccgtc 660 aagcaggcctacgccaccgtcgccaccgcgcggaggtcaagtacactgtctttgag 720 780 accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccqcc 40 gccgctgccaccgccaccgccgccgttggcgcggccaccggcgccaccgcc 840 gctactggtggctacaaagtc 861 Mutant 13: 45 I45K, E133S, P214G, P232G, L243E, Q254K: gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgcc 60 acccccgccgcccggagcggagccagcaggtaaggcgacgaccgaggagcagaag 120 ctgatcgagaagAAAaacgccggcttcaaggcggccttggccgctgccggcggtcccg 180 50 ccagcggacaagtacaggacgttcgtcgcaaccttcggcgcggcctccaacaaggccttc 240 300 gcggagggcctctcgggcgagcccaagggcgccgccgaatccagctccaaggccgcgctc 360 acctccaagctcgacgccgcctacaagctcgcctacaagacagccgagggcgcgacgcct gaggccaagtacgacgcctacgtcgccaccgtaagcAGCgcgctccgcatcatcgccggc 420 480 accetegaggtecacgecgteaagceegeggeegaggaggteaaggteateeeegeegge 55 gagctgcaggtcatcgagaaggtcgacgccgccttcaaggtcgctgccaccgccgccaac 540

gccgccccgccaacgacaagttcaccgtcttcgaggccgccttcaacgacgccatcaag

gcgagcacgggcgcctacgagagctacaagttcatcGGCgccctggaggccgccgtc 660
aagcaggcctacgccgccaccgtcgccaccggGGCgaggtcaagtacactgtctttgag 720
accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagccgcc 780
gccgctgccaccgccaccgcaaccgccgttggcgggccaccggcgccaccgcc 840

Mutant 14:

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I45K, E133S, K211N, P232G, L243E, Q254K:

10	gccgatctcggttacggccccgccaccccagctgccccggccgccggctacacccccgccaccccgccacccgccgccgccgccgccgc	60 120 180 240
15	gcggagggcctctcgggcgagcccaagggcgcgccgaatccagctccaaggccgcgctcacctccaaggccgccgcctacaagcccgaggcgcgacgcctgaggccaagtacgacgcctacgcaccgtaagcAGCgcgctccgcatcatcgccggcacccgcgggcctagggtcaaggtcatcccgccggc	300 360 420 480
20	gagetgeaggteategagaaggtegaegeegeetteaaggtegetgeeacegeegeeace geegeeceegeeateaag gegageacggeggegeetaegagagetaeAACtteateecegeectggaggeegeetteaag aageaggeetaegeegtegeegte	540 600 660 720
	accgcaGAAaaaaaggccatcaccgccatgtccgaagcaAAAaaggctgccaagcccgcc gccgctgccaccgccaccgcaaccgccgctgttggcgcggccaccggcggccaccgcc gctactggtggctacaaagtc	780 840 861

EXAMPLE 9

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T-cell reactivity of recombinant and mutant Bet v 1:

Purpose:

To investigate an *in vitro* T-cell response to the mutated allergens in terms of proliferation and cytokine production.

Methods:

PBL (Peripheral blood lymphocytes) from allergic patients 40 were used in the following investigation.

Eight bet v 1 specific T-cell lines were established from the PBL with naturally purified bet v 1 in order to sustain the variety of bet v 1 isoforms the T-cells are presented to, as described in a previously published

protocol (26).

Ten PBL and eight T-cell lines were stimulated with birch extract (Bet v), naturally purified bet v 1 (nBet v 1), recombinant Bet v 1 (rBet v 1 or wt; 27) and four different mutated forms of rBet v 1 (described elsewhere): 2595, 2628, 2637, 2744, 2773. The 2637 mutant was later found to be partly unfolded and will not be discussed.

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In brief: In a round-bottomed 96 well plate PBL were added in 2 x 10^5 per well. The different birch samples were added in three different concentrations in quadroplicates and allowed to grow for 6 days. At day 6 cell half of volume (100 μ l) from each well with the highest concentration of birch were harvested for cytokine production. Radioactive labelled thymidine was added to the wells. Next day (day 7) the cells were harvested on a filter. Scintilation fluid was added to the filter and the radioactivity was measured in a scintilation counter.

Likewise in a 96 well round-bottomed 96 well plate T-cells were added in 3×10^4 T-cells per well and stimulated with irradiated autologous PBL (1×10^5 cells/well) and 3 different concentrations of the different birch samples. After 1 day cells from each well with the highest concentration birch were harvested for cytokine production. Radioactive labelled thymidine were added to the wells. At day 2 the cells were harvested onto a filter and counted as described for PBL.

Supernatant from the quadroplicates were pooled and cytokines were measured using a CBA (cytokine bead array) kit from Becton Dickinson.

Results:

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Ten PBL cultures showed specific stimulation to birch. In general proliferation of the PBL to the different birch samples were similar, although variations could be seen. In 3 PBL, nBet v 1 stimulated proliferation better than rBet v 1 and the mutants. The mutant birch samples stimulated PBL almost identical to rBet v 1 (Fig. 41). Fig. 41 shows the Stimulation Index for the abovementioned Bet v 1 preparations. The Stimulation Index (SI) is calculated as proliferation (cpm: count per minute) of the stimulated sample (highest concentration) divided with the proliferation (cpm) of the medium control. PPD designates purified protein derivative from mucobacterium tuberculosis, which serves as a positive control.

Cytokine production was dominated by IFN-gamma and increased proportionally with PBL proliferation. No signs of a Th1/Th2 shift were apparent (Fig. 42-44). Figure 42 shows a patient with a Th0 profile, Figure 43 a Th1 profile and Figure 44 a Th2 profile. Cytokine production is measured in pg/ml indicated as the bars and the ratio between IL-5/IFN-gamma is the lower dashed line (Y-axis to the right). Proliferation is measured in cpm seen on the Y-axis to the right as a solid line measured in cpm. Medium and MBP (maltose bindig protein) are included as background controls.

30 Eight T-cell lines established on nBet v 1 and all, except one, proliferated equally well to all birch samples. Four T-cell lines were secreting ThO like cytokines based on the IL-5 and IFN-gamma ratio (Th2 > 5, 5 > ThO > 0.2, 0.2 > Th1). Three T-cell lines were secreting Th1 cytokines and one T-cell line was secreting Th2 cytokines. The IL-5/IFN-gamma ratio was not affected

by the different birch samples.

Conclusion:

5 All PBL cultures and 7/8 T-cell lines that showed specific stimulation to nBet v 1 did also respond to rBet v 1 and the mutants. These data suggests that for T-cell stimulation a single isoform of Bet v 1 or these 4 mutants can substitute for the mixture of individual isoforms found in the natural allergen preparations. Thus, vaccines based on recombinant allergens or these 4 mutants will address the existing Bet v 1 specific T-cell population.

15 EXAMPLE 10

Induction of Bet v 1 specific IgG antibodies and blocking antibodies following immunization with recombinant and mutant Bet v 1 proteins:

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In this section the term "blocking antibodies" is defined as antibodies, different from human IgE antibodies, that are able to bind to an antigen and prevent the binding of human IgE antibodies to that antigen.

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The ability of recombinant Bet v1 2227 wild type protein (rBet v 1) and Bet v 1 2595, 2628, 2744 and 2773 mutant proteins to induce Bet v 1 specific IgG antibodies and blocking antibodies was tested in immunization experiments in mice.

BALB/cA mice (8 in each group) were immunized by intraperitoneal injections with recombinant Bet v1 2227 wild type protein or the four mutant proteins. The mice were immunized four times with a dose interval of 14 days. The different proteins were conjugated to 1,25

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mg/ml Alhydrogel, (Aluminium Hydroxide gel, 1,3 % pH 8.0 - 8.4, Superfos Biosector). The mice were immunized with either 1 ug protein/dose or 10 ug protein/dose. Blood samples were drawn by orbital bleed at day 0,14,35, 21, 49 and 63.

Specific IgG antibody levels was analyzed by direct ELISA using rBet v 1 coated microtiterplates and biotinylated rabbit anti mouse IgG antibodies (Jackson) as detection antibody. Immunization with recombinant Bet v1 2227 wild type protein or the four mutant proteins induced a strong r Bet v 1 specific IgG response. This finding demonstrates that the four mutated proteins are able to induce antibodies that are highly cross reactive to the Bet v 1 2227 wild type protein

To assess the induction of blocking antibodies, serum samples birch pollen allergic patients from incubated with paramagnetic beads coated with monoclonal mouse anti-human IgE antibody. incubation, the beads were washed and resuspended in buffer or diluted samples (1:100) of mouse serum from unimmunized mice (control) or mice immunized as described above. Biotinylated r Bet v 1 was then added to this mixture of beads and mouse serum antibodies. After incubation, the beads were washed and bound biotinylated detected 1 was using acridinium streptavidine. Incubation of beads with serum from unimmunized mice did not change the binding of r Bet v 1 to the beads. In contrast, incubation of the beads with serum from mice immunized with the recombinant Bet v1 2227 wild type protein or the four mutant proteins significantly reduced binding of r Bet v 1 to the beads demonstrating the presence of Bet v 1 specific blocking antibodies in the serum samples. Thus, at day 63 one or more serum samples from all high dose (10 ug/dose)

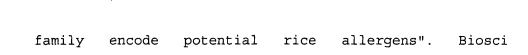
immunization groups were able to reduce binding of r Bet v1 to the beads with more than 80%. These findings demonstrate that the four mutated proteins are able to induce antibodies that can act as Bet v 1 specific blocking antibodies.

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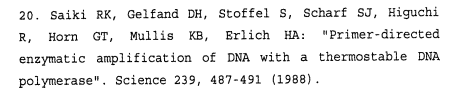


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